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Suresh et al.

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(54) **SYRINGE ASSEMBLY HAVING DISABLING MECHANISM**

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(*) Notice: Subject to any disclaimer, the term of this patent is extended or adjusted under 35 U.S.C. 154(b) by 631 days.

This patent is subject to a terminal disclaimer.

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Primary Examiner—Matthew DeSanto

(57) **ABSTRACT**

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A61M 5/00 (2006.01)
A61M 5/315 (2006.01)

(52) **U.S. Cl.** **604/110**; 604/228

(58) **Field of Classification Search** 604/110, 604/181, 187, 198, 218, 223, 228, 241, 227
See application file for complete search history.

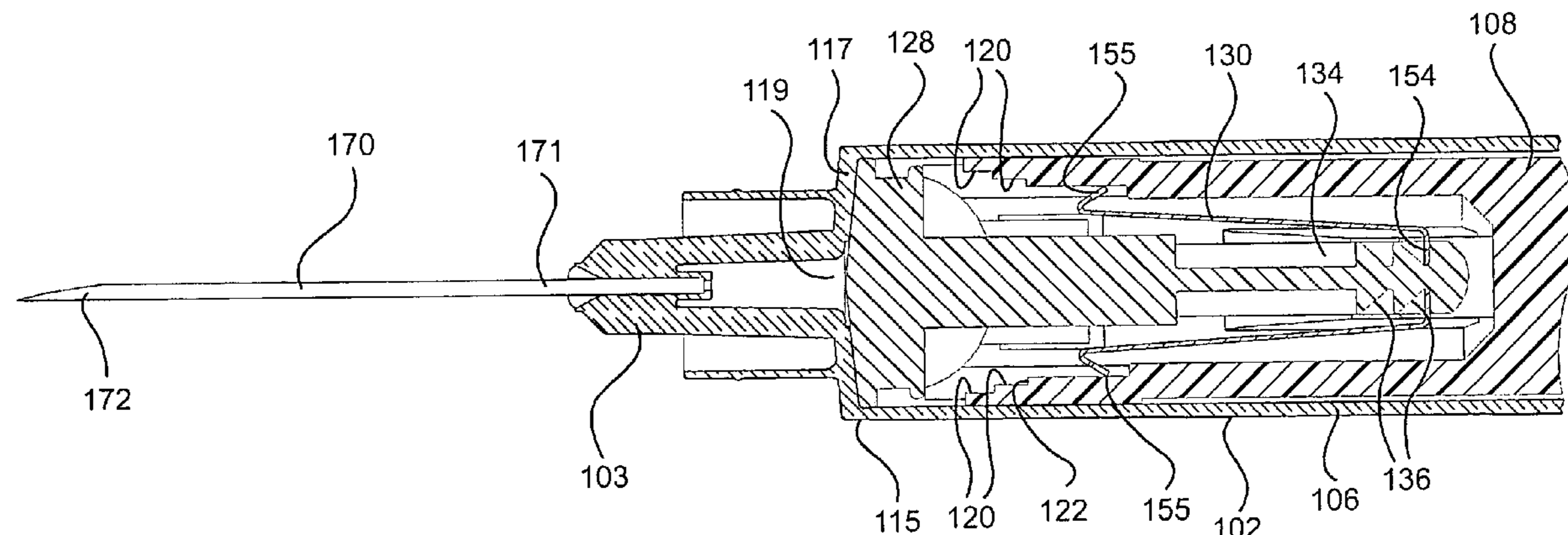
A syringe assembly having passive disabling structure includes a barrel and a plunger rod assembly. The plunger rod assembly includes a plunger rod and a stopper connected by an indexing locking element. The number of strokes of the syringe plunger before the stopper is locked into the barrel rendering the syringe assembly unusable is determined by the number of detents on the plunger rod and stopper which engage the locking mechanism. Upon completion of the final delivery stroke, any attempt to withdraw the plunger rod from the barrel will cause the locking element to engage the barrel and trap the stopper in the barrel preventing further use of the syringe.

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27 Claims, 24 Drawing Sheets



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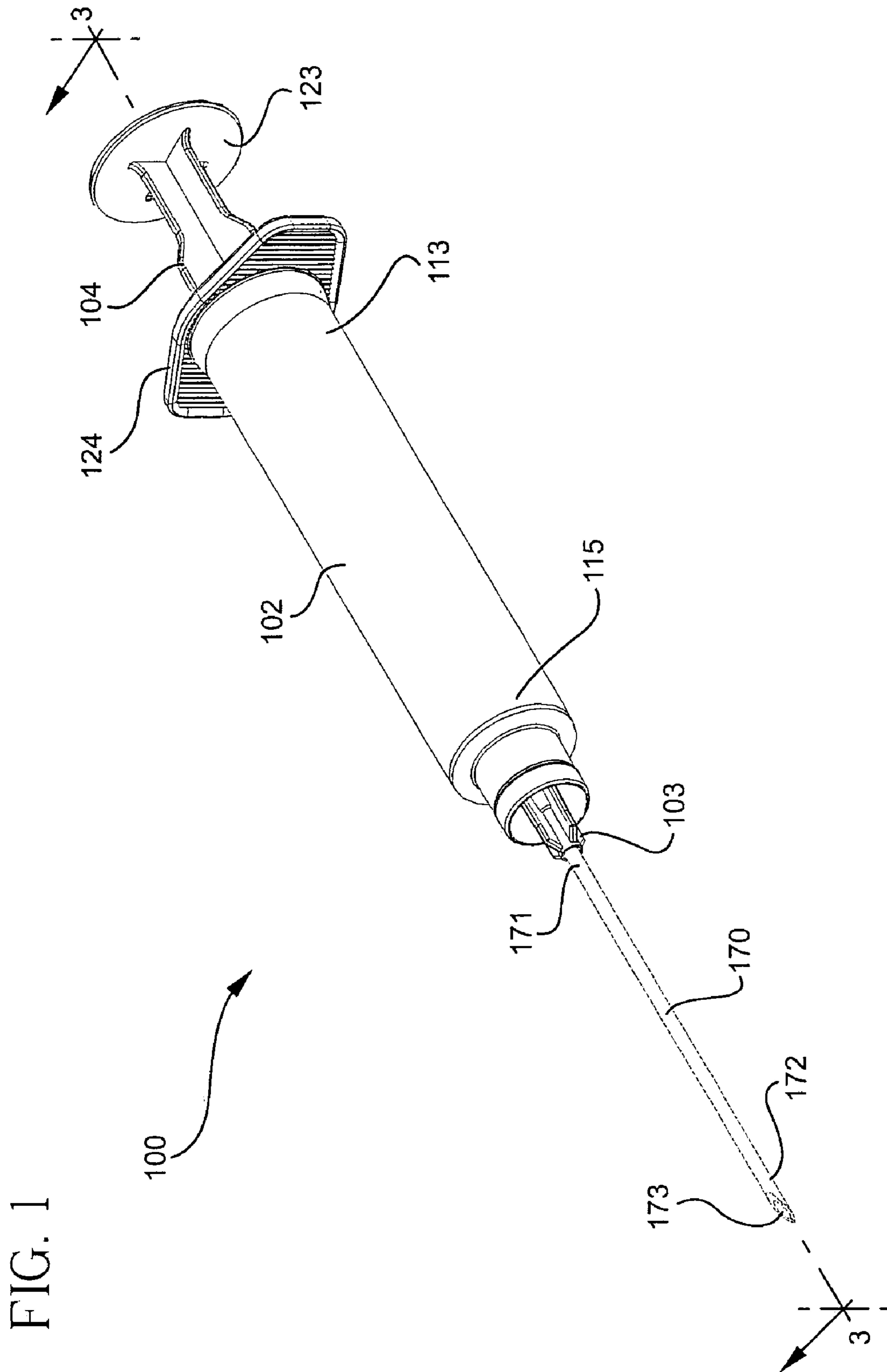
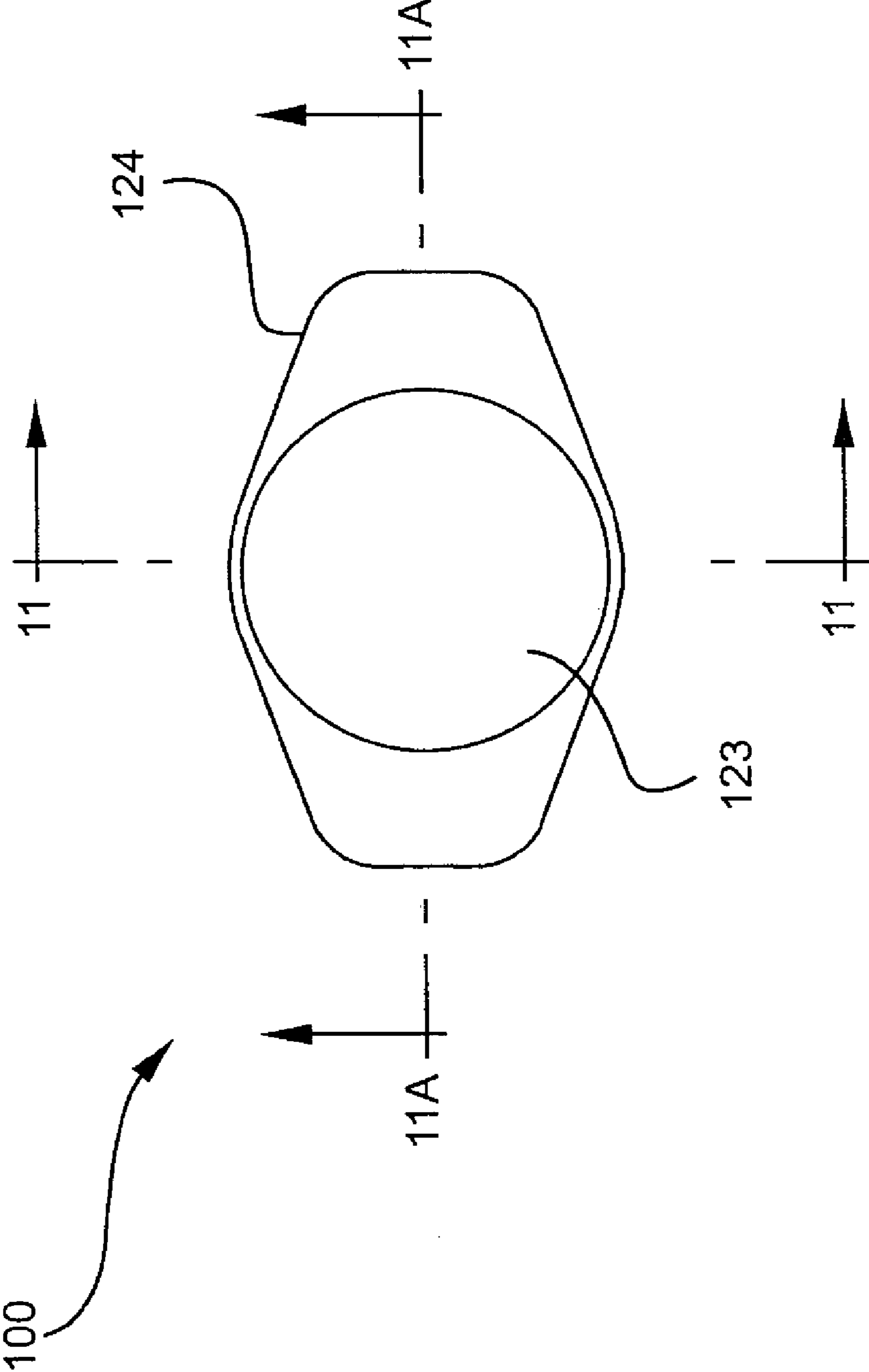


FIG. 2



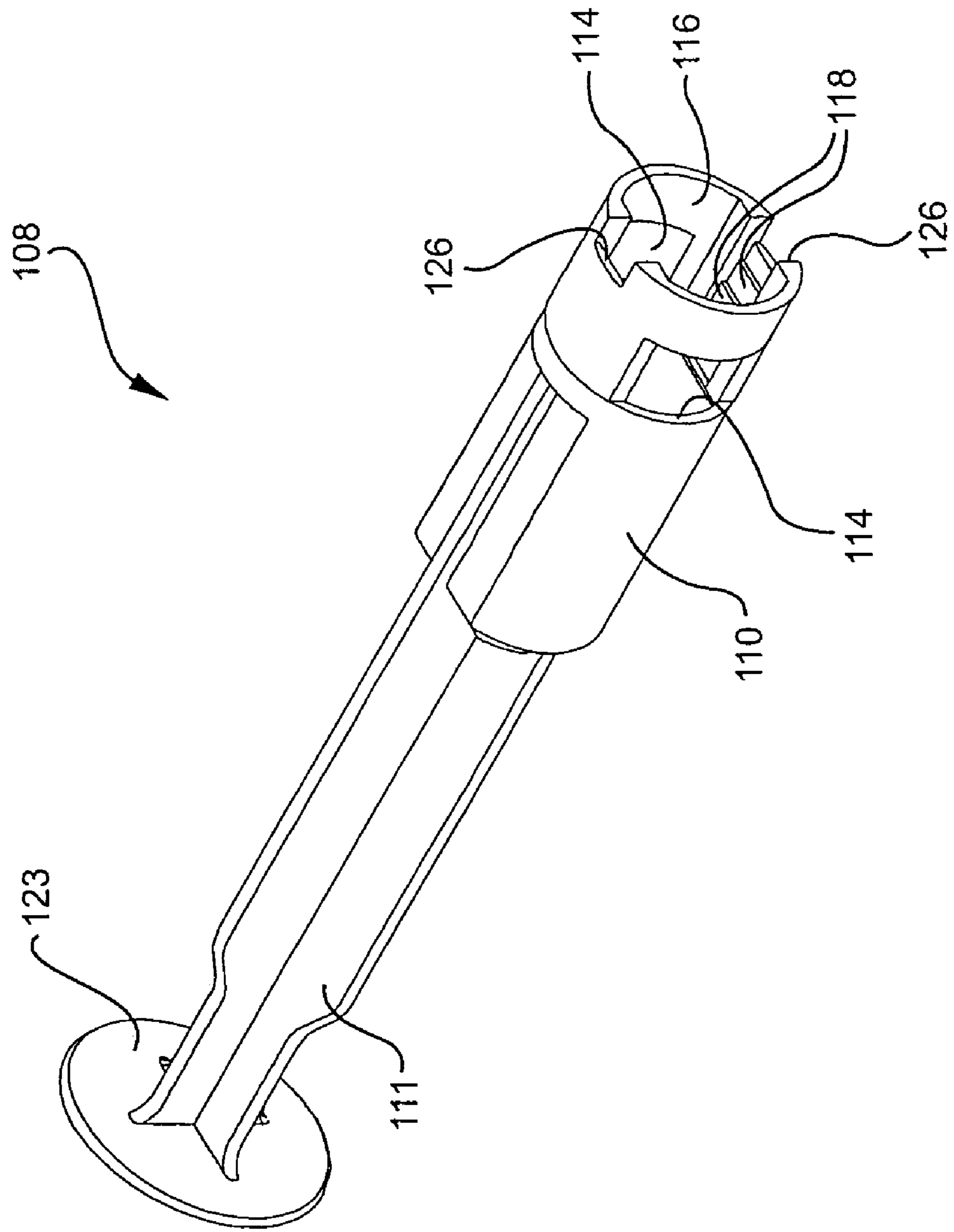


FIG. 4

FIG. 5

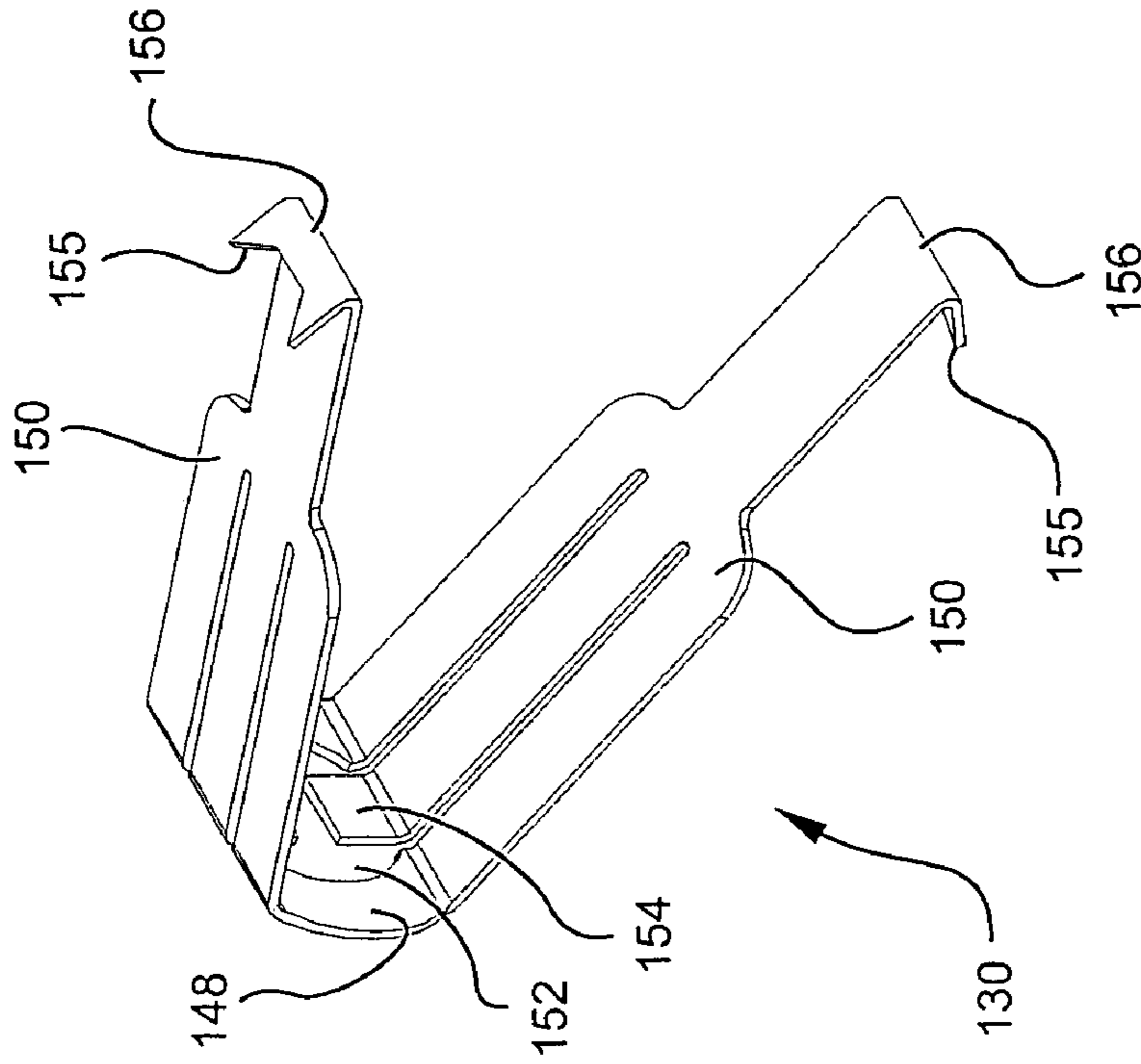
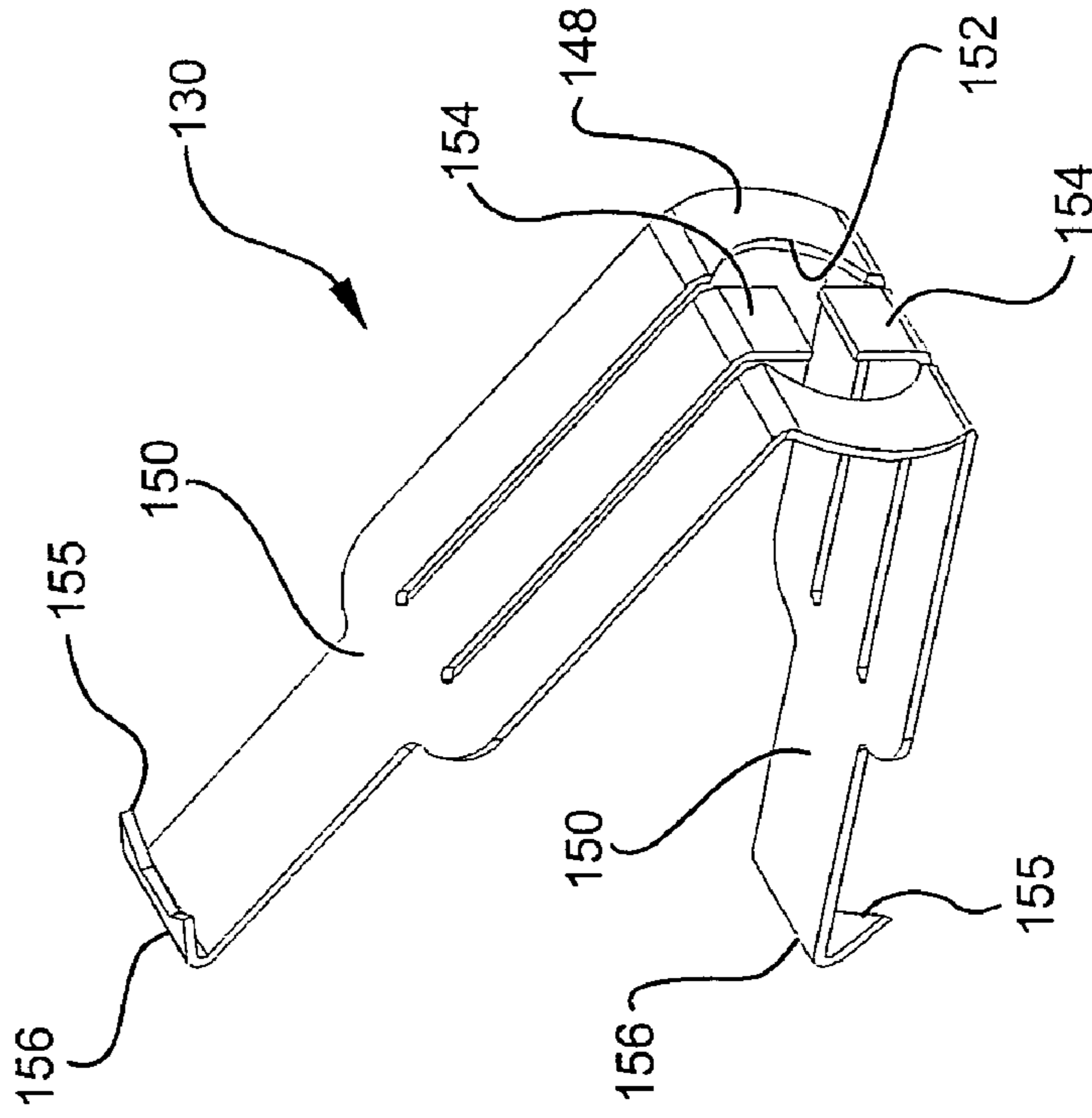


FIG. 6



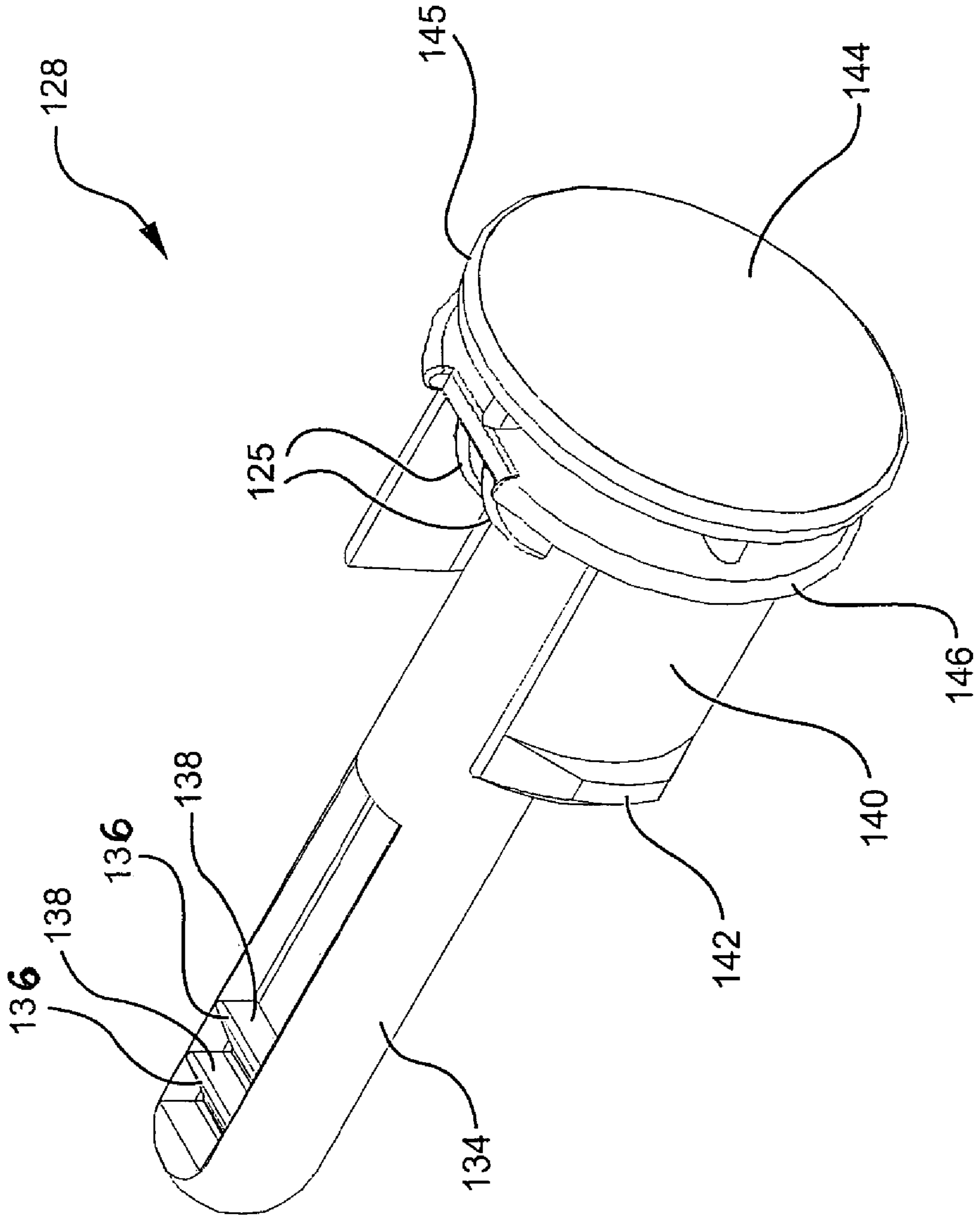


FIG. 7

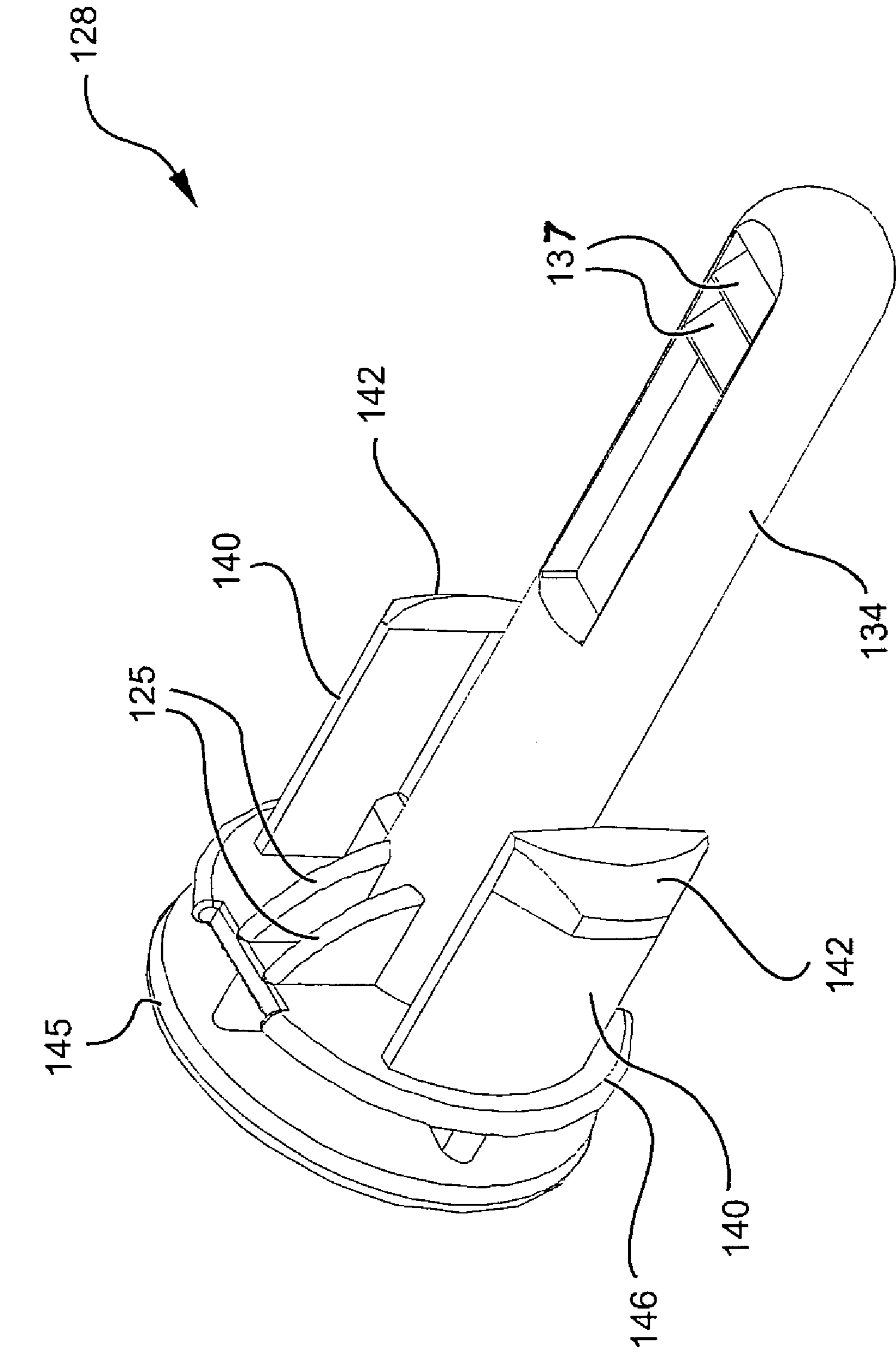


FIG. 8

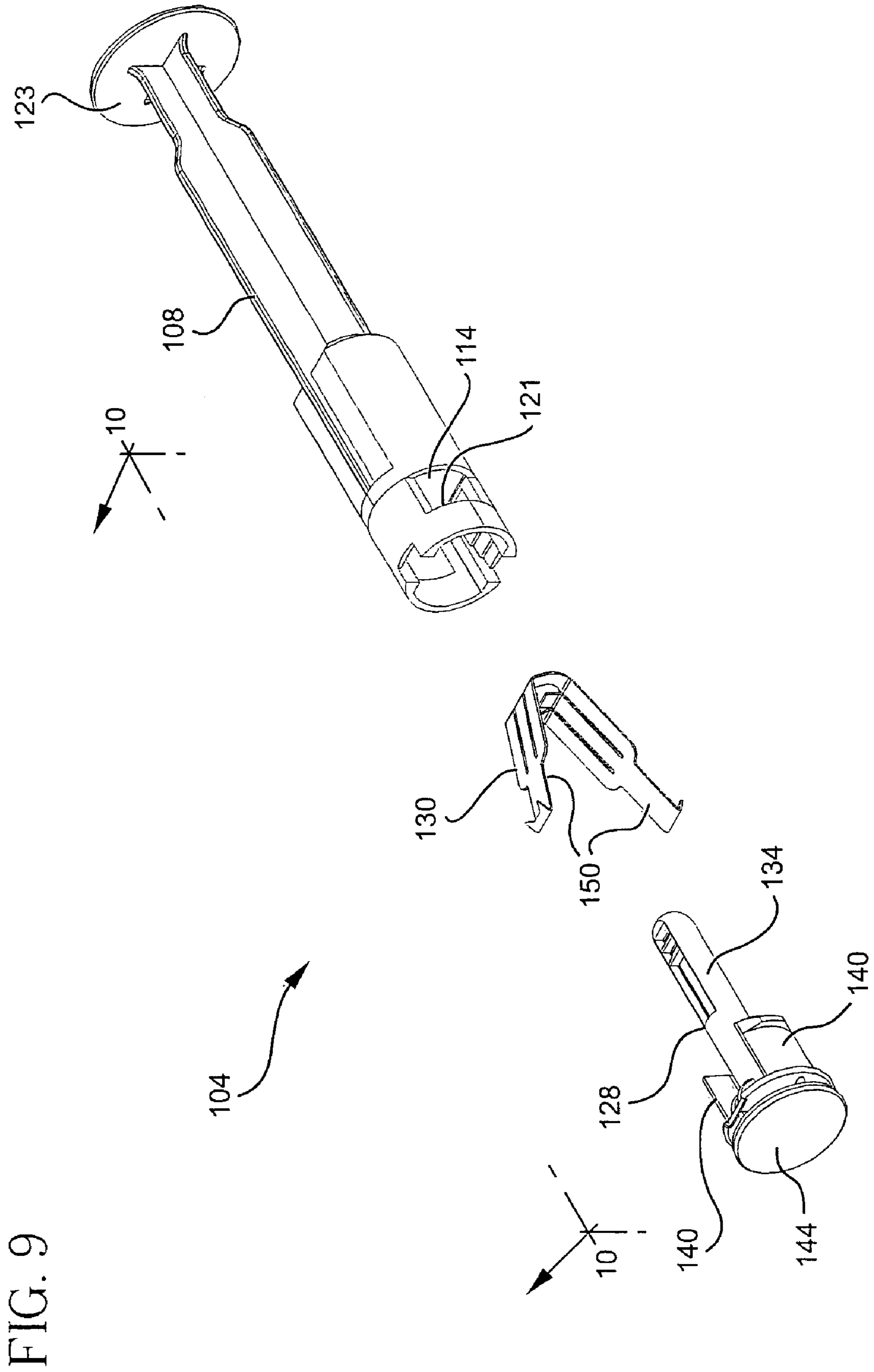


FIG. 10

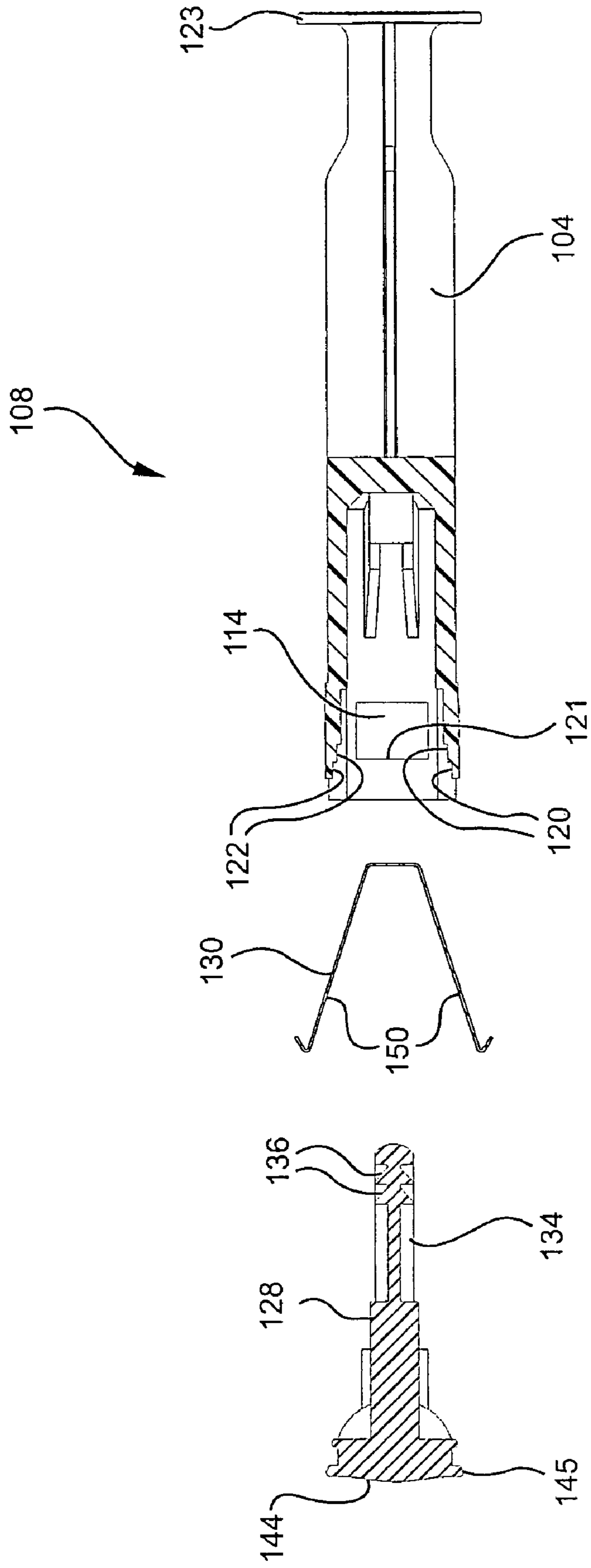


FIG. 11

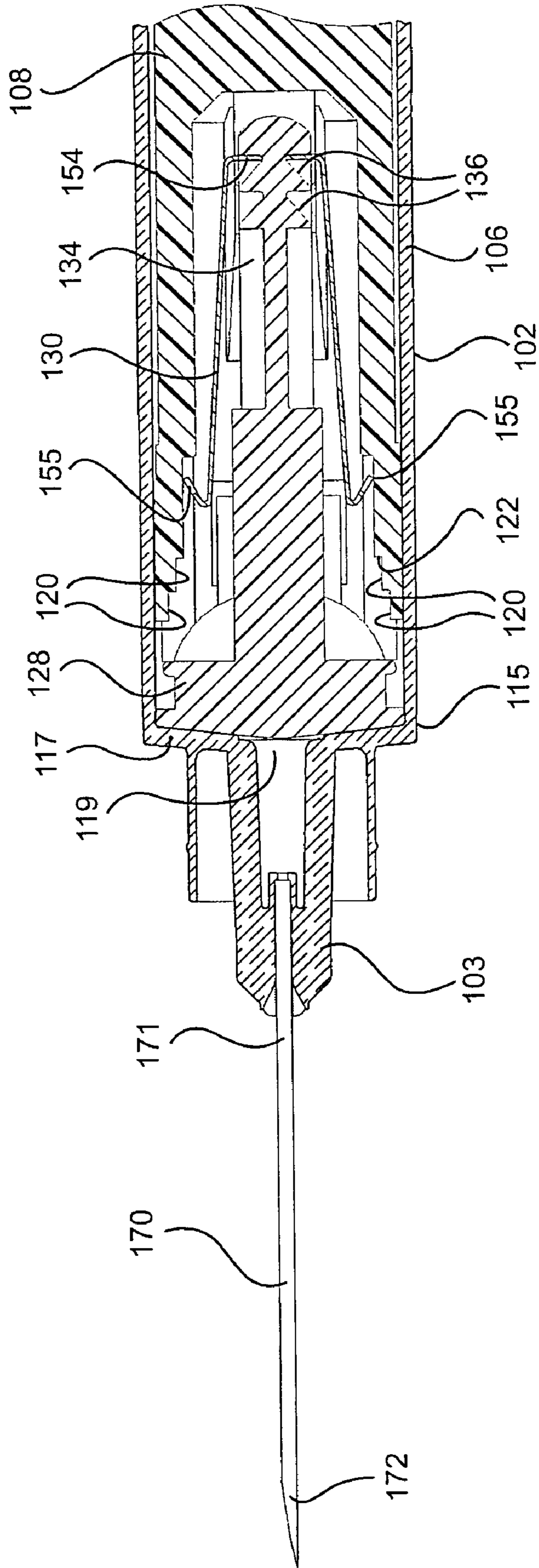


FIG. 11A

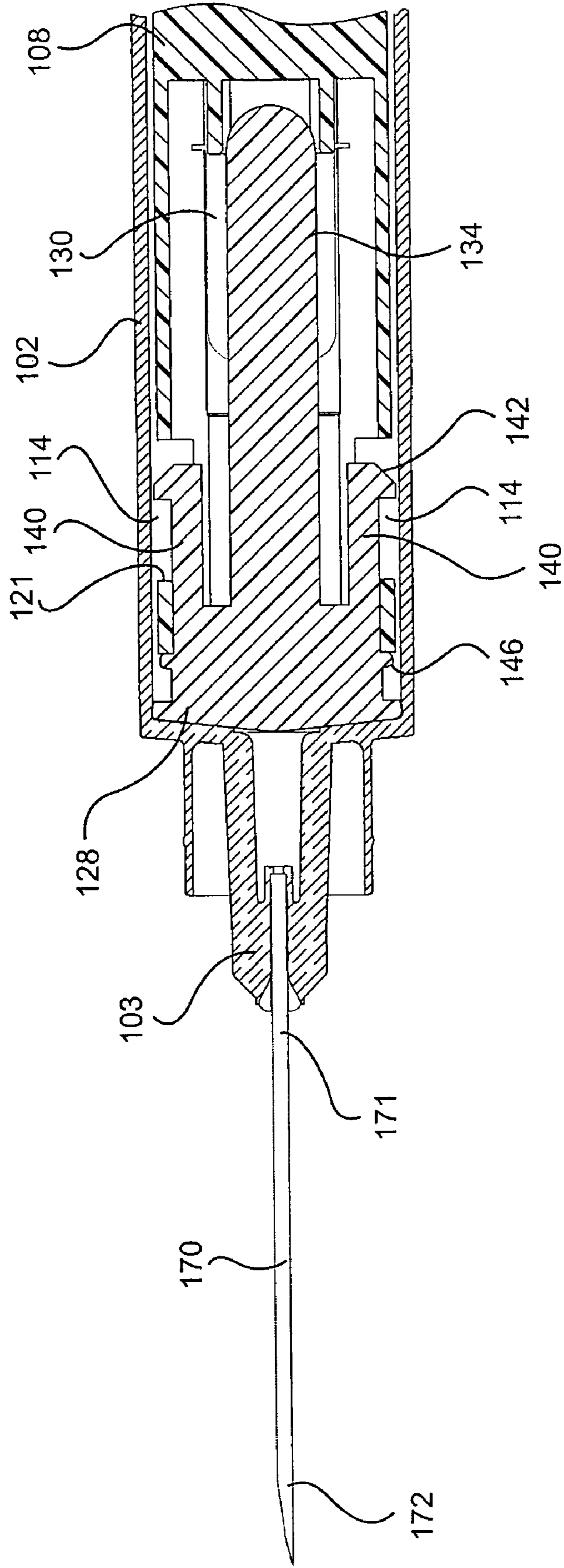


FIG. 12

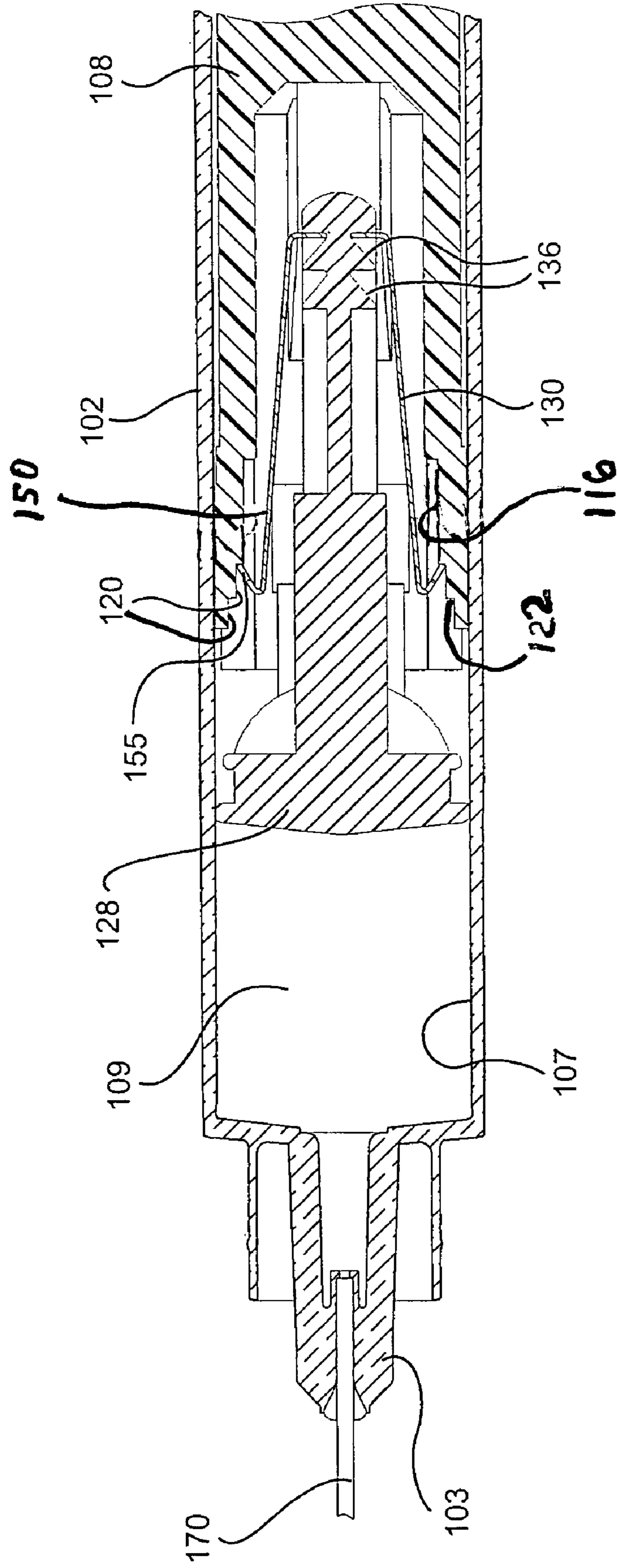


FIG. 12A

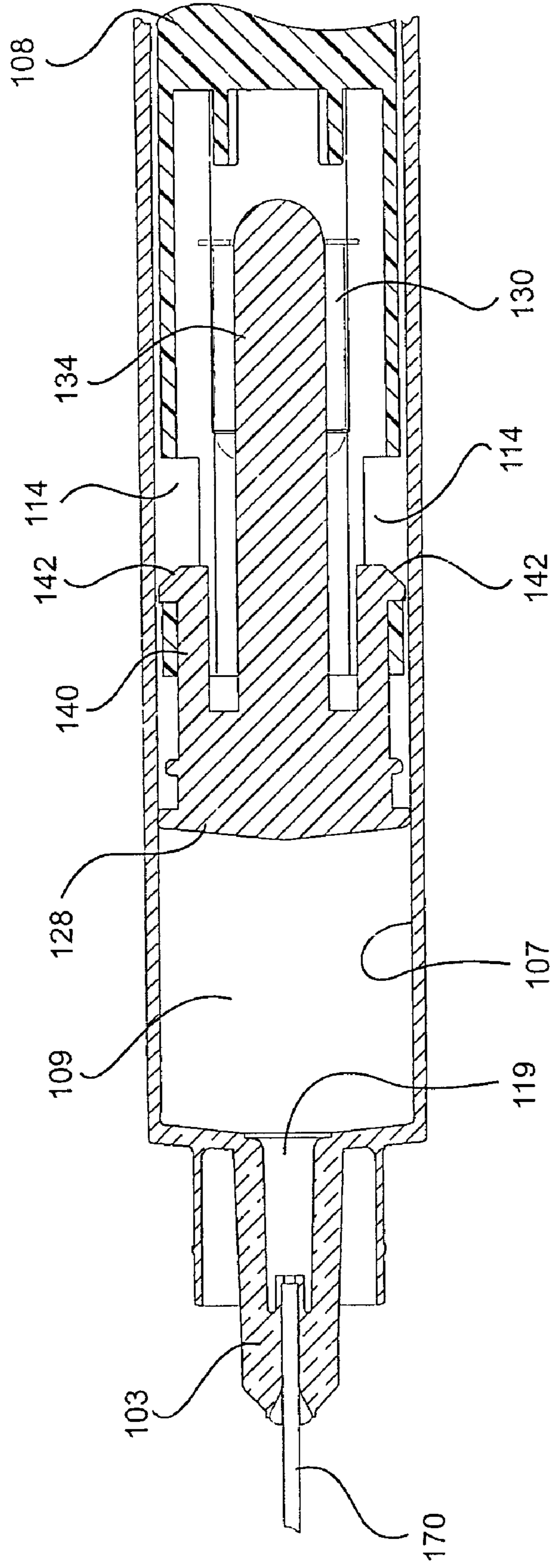


FIG. 13

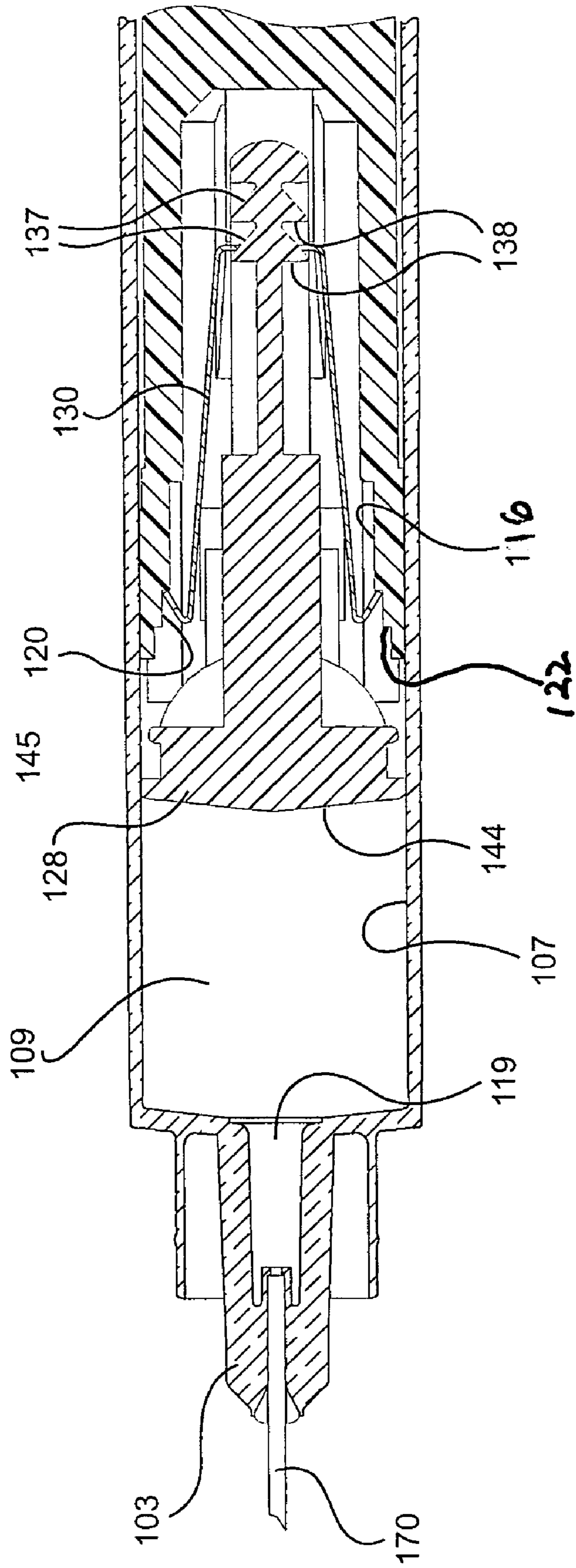


FIG. 13A

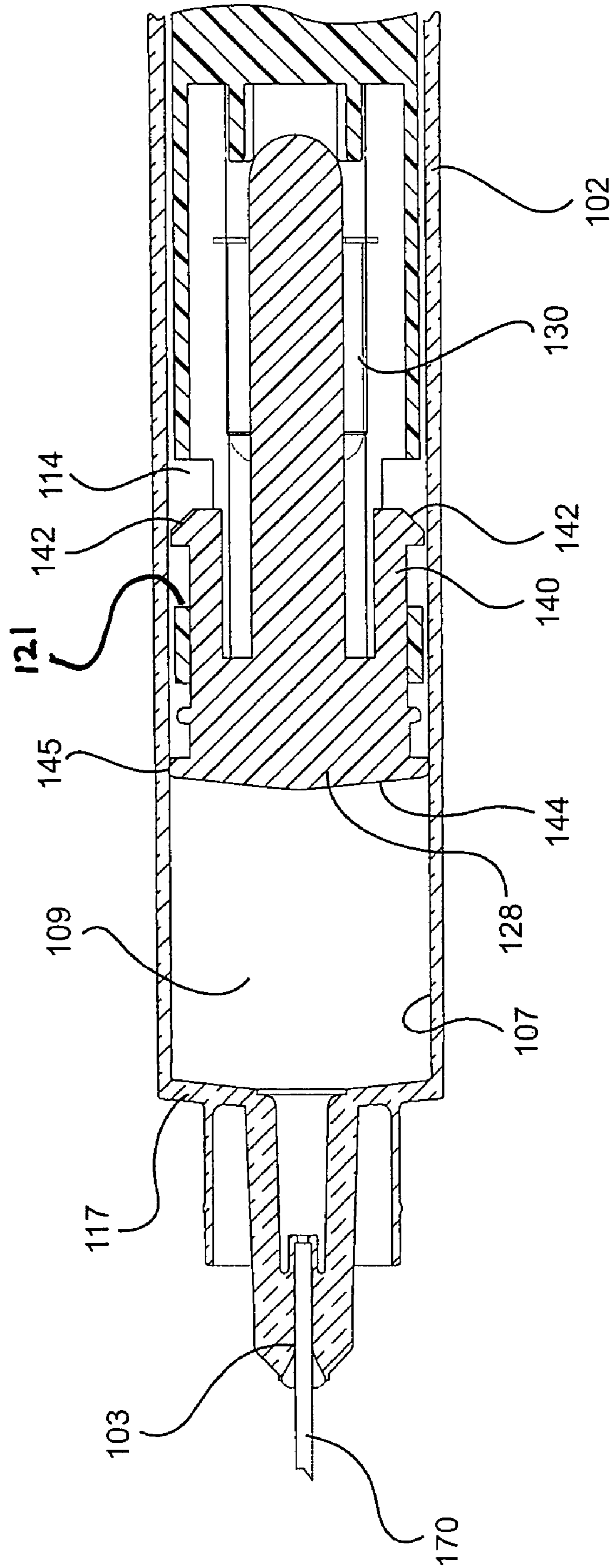


FIG. 14

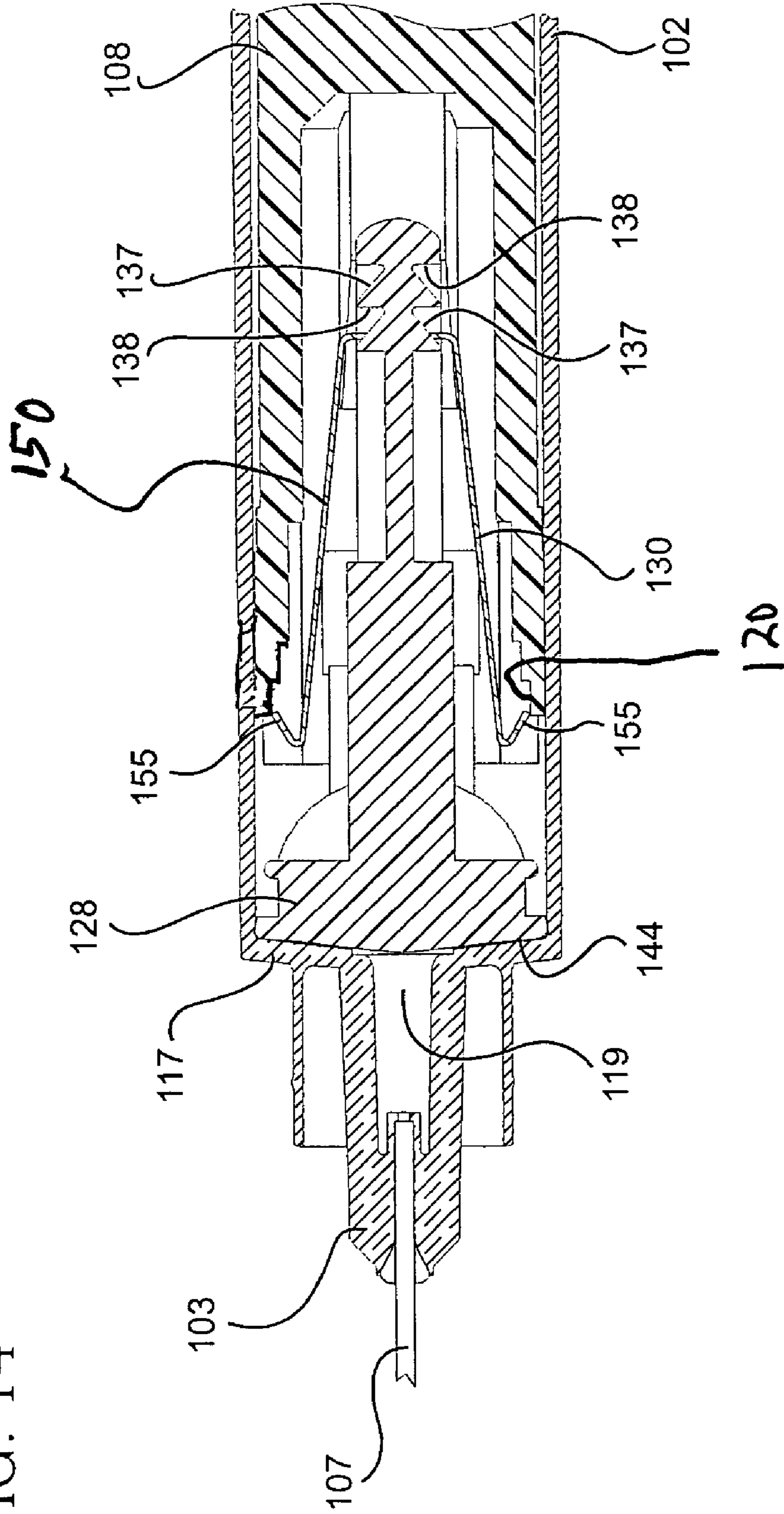


FIG. 14A

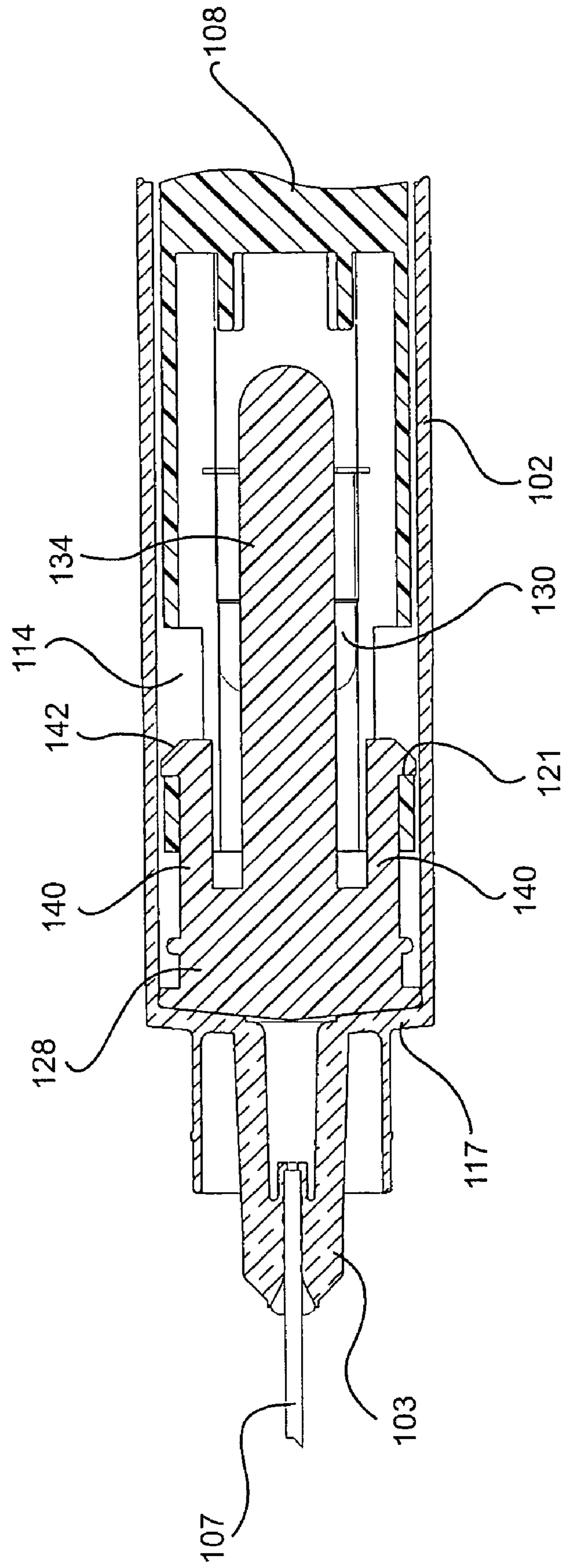


FIG. 15

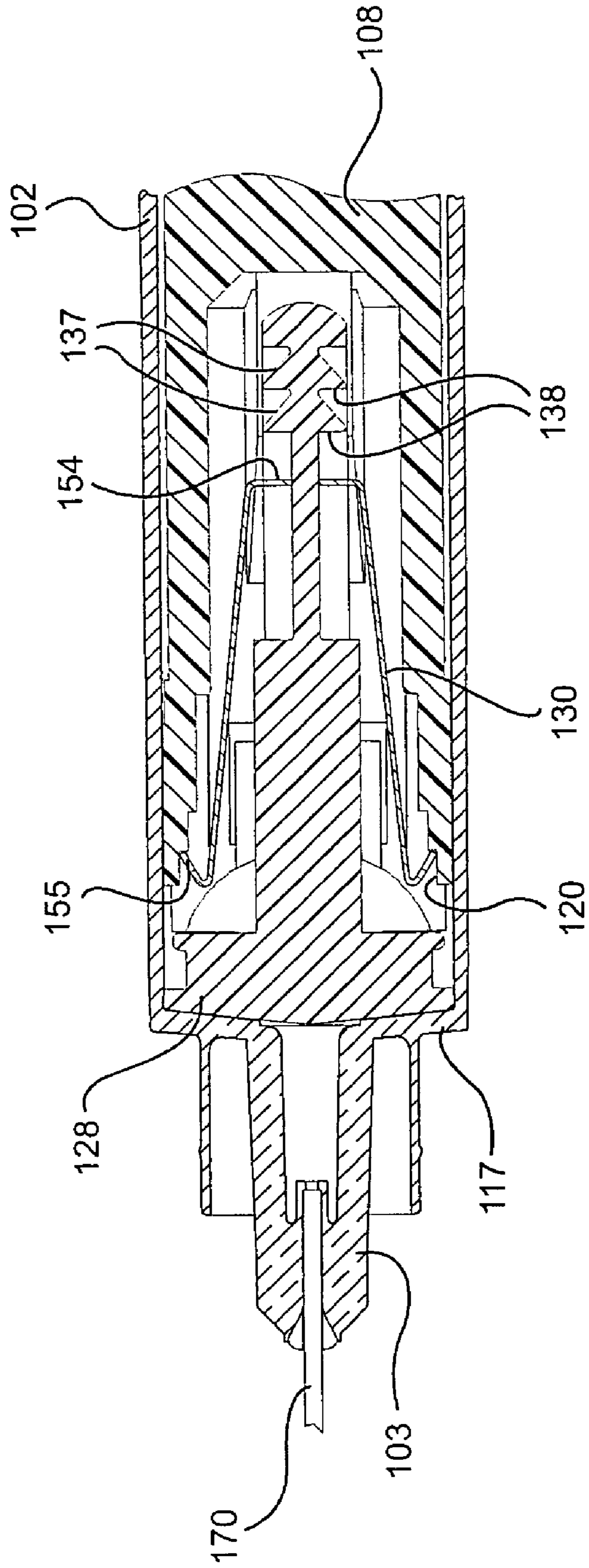


FIG. 15A

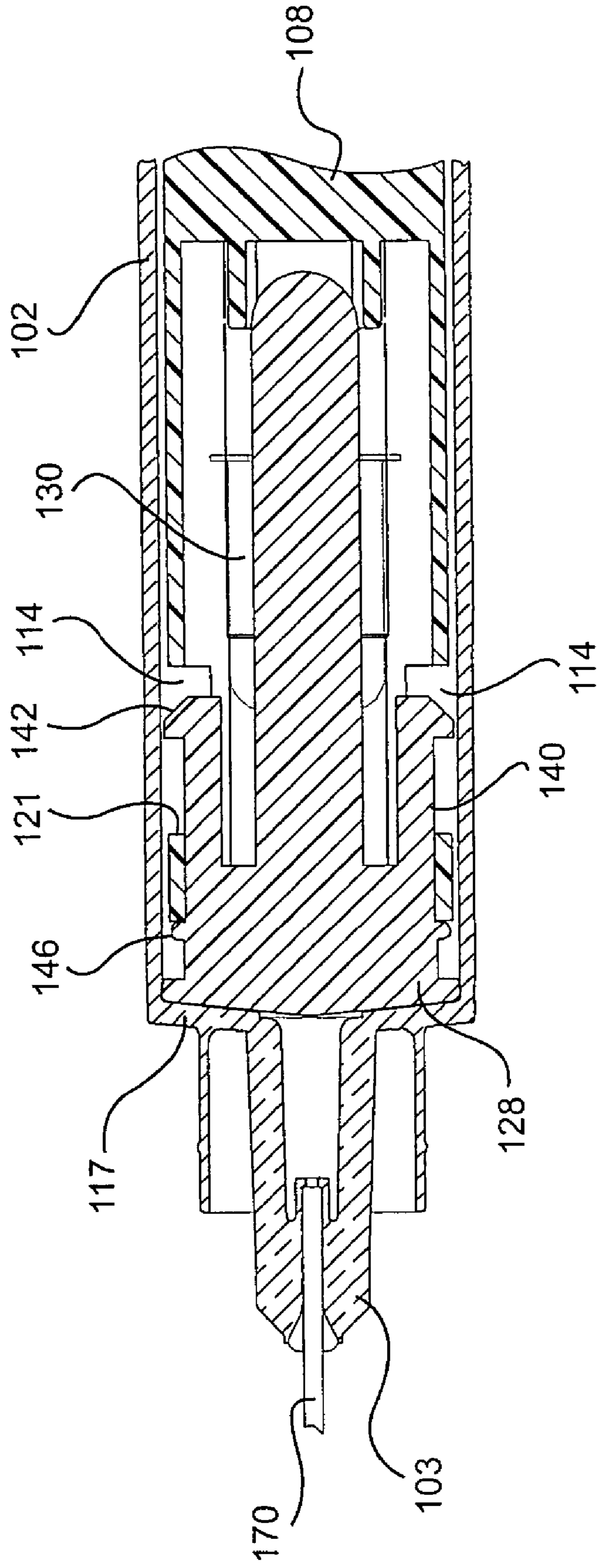


FIG. 16

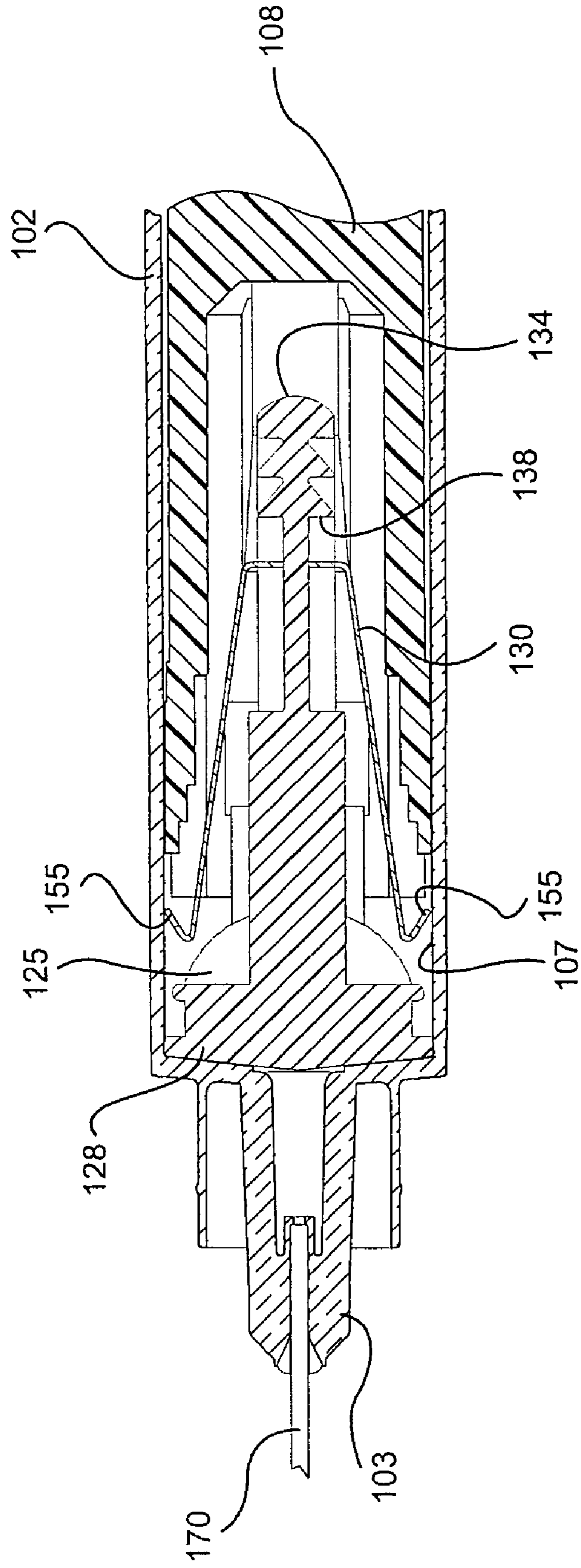


FIG. 16A

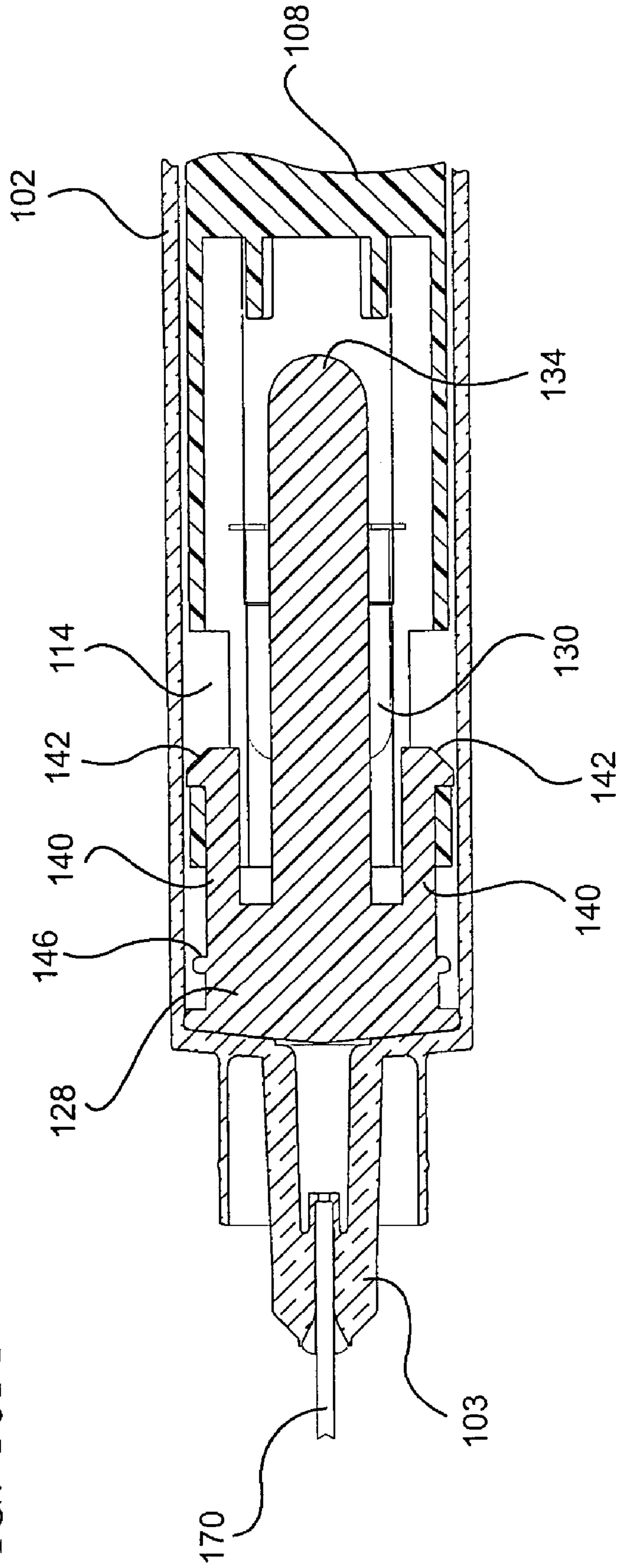
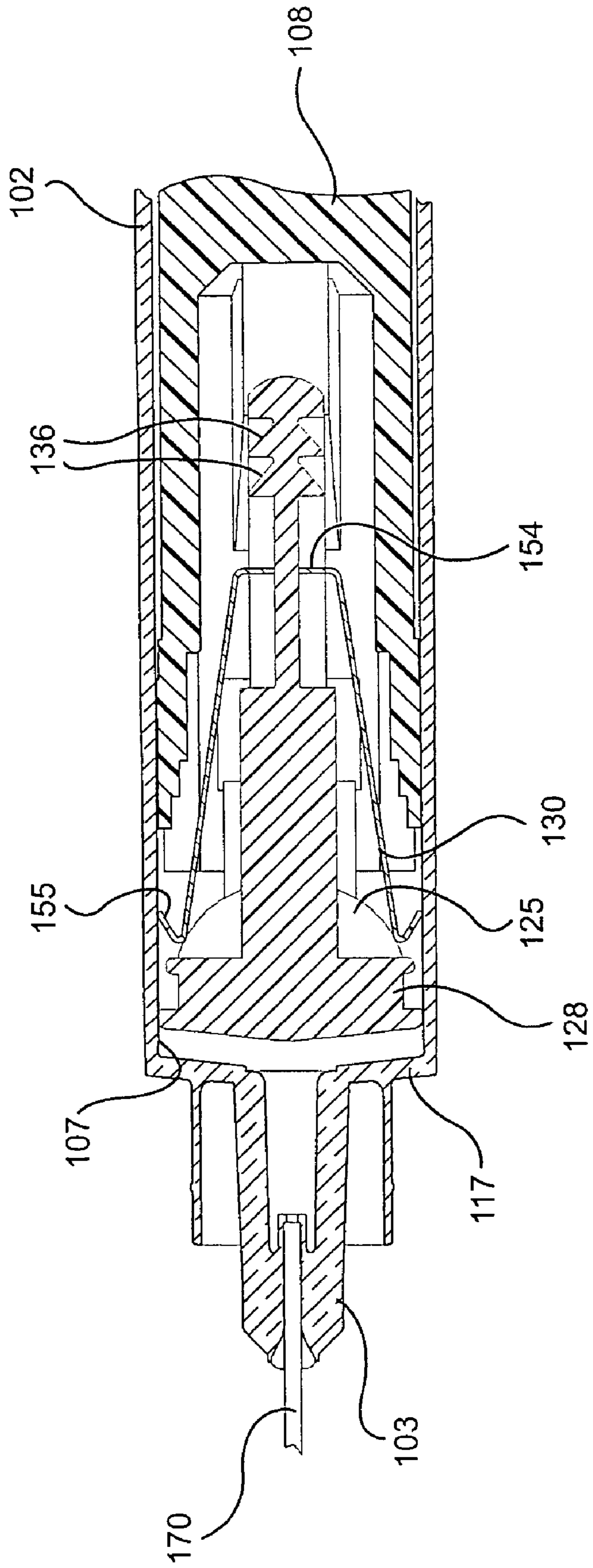
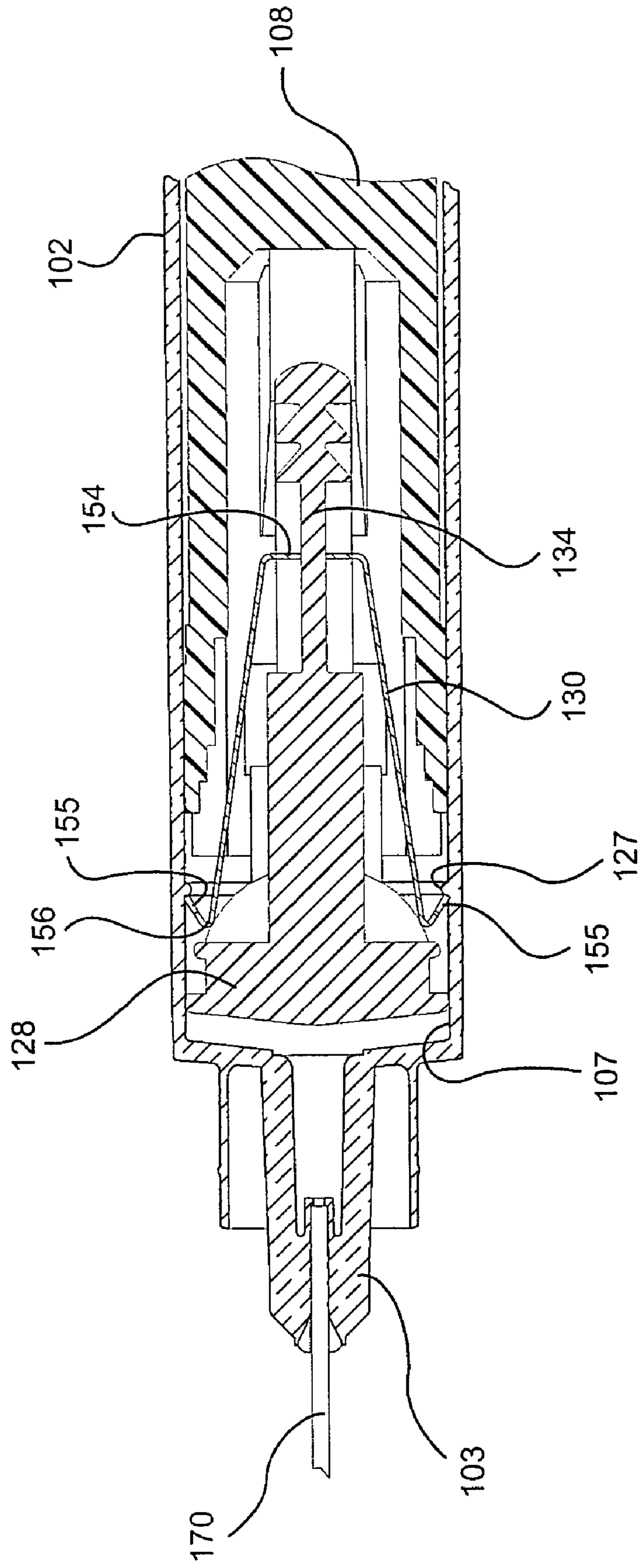


FIG. 17





SYRINGE ASSEMBLY HAVING DISABLING MECHANISM

This application claims priority from U.S. Provisional Application No. 60/490,939, filed Jul. 30, 2003 and U.S. Provisional Application No. 60/523,490, filed Nov. 20, 2003.

BACKGROUND OF THE INVENTION

The present invention relates to syringe assemblies and particularly to syringe assemblies having an automatic disabling mechanism.

Throughout the world the multiple use of hypodermic syringe products which are intended for single-use only, is instrumental in drug abuse and in the transfer of contagious diseases. Intravenous drug users who routinely share and re-use syringes are a high-risk group with respect to the AIDS virus. Also, the effects of multiple use are a major concern in some countries where repeated use of syringe products during mass immunization programs may be responsible for the spread of many diseases. Re-use of single-use hypodermic syringe assemblies is also instrumental in the spread of drug abuse even in the absence of infection or disease.

Many attempts have been made to remedy this problem. Most notable are early contributions which relied on a specific act to destroy the syringe after use either by using a destructive device or providing a syringe assembly with frangible zones so that the syringe could be rendered inoperable by the application of force. Other attempts involve the inclusion of structure which would allow the destruction or defeating of the syringe function to a conscious act by the syringe user. Although many of these devices work quite well, they do require the specific intent of the user followed by the actual act to destroy or render the syringe inoperable. These devices are not effective with a user having the specific intent to re-use the hypodermic syringe. Accordingly, there was a need for a single-use hypodermic syringe which after use will become inoperable or incapable of further use automatically without any additional act on the part of the user. The automatic function is much harder to provide because the means for rendering the syringe inoperable must not prevent its filling or use under normal conditions.

A single-use syringe which automatically disables after injection is taught in U.S. Pat. No. 4,973,310 Kosinski. This syringe contains a locking element positioned in the syringe barrel between the plunger rod and the inside surface of the barrel. In use, the syringe allows the user to draw a pre-selected amount of medication into the chamber of the barrel and deliver this medication, as through injection, into the patient. Any attempt to withdraw the plunger to use the syringe a second time will cause the locking element to embed itself into the inside surface of the syringe barrel to prevent proximal motion of the plunger rod.

There is still a need for a single-use syringe which will allow a pre-selected number of plunger rod strokes before the automatic disabling mechanism activates. For example, four strokes of the plunger may be required to complete the injection process. Such as when the syringe assembly is used to draw a diluent into the syringe barrel, dispense the diluent into a vial containing the substance to be reconstituted, drawing back the reconstituted medication into the syringe and then delivering the contents of the syringe into the patient.

SUMMARY OF THE INVENTION

An operable syringe assembly having a passive disabling structure includes a barrel having a cylindrical sidewall with an inside surface defining a chamber for retaining fluid, an open proximal end and a distal end including a distal wall having a passageway therethrough in fluid communication with the chamber. An elongate hollow plunger rod having a proximal end, an open distal end, and an interior surface is provided. A stopper has a circular-shaped sealing element having a peripheral surface forming a seal with the inside surface of the barrel and a boss member projecting proximally from the sealing element. A locking element interacts between the stopper and the plunger rod. The locking element includes a central body portion having at least one cantilevered leg extending distally outwardly from the body portion wherein the leg includes a sharp free end directed outwardly for engaging the inside surface of the barrel. The locking element is movably connected to the boss of the stopper and movably connected to the plunger rod interior surface. Structure for indexing the locking element distally in the plunger rod during proximal motion of the plunger rod to draw fluid into the chamber and for indexing the locking element distally on the boss of the stopper during distally-directed motion of the plunger rod for delivering fluid from the chamber through the passageway and means for engaging the locking element with the inside surface of the barrel sidewall for preventing reuse of the syringe assembly is provided.

Structure or means for engaging the locking element with the inside surface of the barrel sidewall may include an opening in the distal end of the plunger rod to shorten the axial length of the interior surface area of the opening for allowing the sharp free end to project outwardly past the plunger rod and onto the inside surface of the barrel.

The structure or means for indexing includes at least one detent on the interior surface of the distal end of the plunger rod, at least one boss detent on the boss and at least one cantilevered arm on the stopper having an outwardly extending rib near its free end sized to engage a recess in the inside surface of the plunger rod. The indexing structure further includes the locking element having at least one finger element extending inwardly from an aperture in the central body portion of the locking element. The locking element is positioned with its sharp free end contacting the interior surface of the plunger rod proximally of the at least one detent and the boss member is positioned in the aperture of the locking element wherein the at least one finger element is contacting the boss member proximally of the at least one boss detent and the outwardly extending rib is positioned in the recess of the plunger rod.

The syringe assembly may be configured so that the at least one detent in the plunger rod includes two axially-spaced detents and the at least one detent in the boss includes two axially-spaced boss detents so that the plunger rod can be moved distally one or two times before proximal motion of the plunger rod causes the locking element to engage the inside surface of the barrel, depending on the initial position of the locking element at the time of use. The two axially-spaced detents in the plunger rod may include two axially-spaced steps each having a blunt surface at its distal end extending inwardly from the interior surface of the plunger rod. The two axially-spaced boss detents may each include an inclined surface extending proximally inwardly and a blunt surface at a distal end of each of the inclined surfaces extending radially inwardly.

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The at least one cantilevered leg of the locking element may include two cantilevered legs positioned on opposite sides of the central body portion. The stopper may further include two radial cam projections positioned to contact and force the two cantilevered legs outwardly when excessive proximally-directed force is applied to the plunger rod in an attempt to overcome the locking element's engagement to the inside surface of the barrel.

The at least one cantilevered arm of the stopper may include two cantilever arms positioned on opposite sides of the boss and said at least one recess in the inner surface of the plunger rod may include two recesses positioned on opposite sides of the plunger rod and configured to receive the outwardly extending ribs of the two cantilevered arms.

The syringe barrel may further include an elongate tip extending distally from the distal wall and having a passageway therethrough in fluid communication with the chamber of the syringe barrel. The syringe assembly may also include a needle cannula having a distal end, a proximal end and a lumen therethrough, wherein the proximal end of the needle cannula is connected to the distal end of the syringe barrel so that the lumen is in fluid communication with the passageway of the barrel. The syringe assembly may include a locking element made of sheet metal such as stainless steel. Further, the stopper and all its elements may be integrally formed of thermoplastic material.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 is a side-elevational view of the syringe assembly of the present invention.

FIG. 2 is a side-elevational end view of the proximal end of the syringe assembly of FIG. 1.

FIG. 3 is a cross-sectional view of the syringe assembly of FIG. 1 taken along line 3-3.

FIG. 4 is a perspective view of the plunger rod of the syringe assembly viewed from its distal end.

FIG. 5 is a perspective view of the locking element of the syringe assembly viewed from its distal end.

FIG. 6 is a perspective view of the locking clip viewed from its proximal end.

FIG. 7 is a perspective view of the stopper of the syringe assembly viewed from its distal end.

FIG. 8 is a perspective view of the stopper viewed from its proximal end.

FIG. 9 is an exploded side-elevational view of the plunger assembly.

FIG. 10 is a cross-sectional view of the plunger assembly of FIG. 9 taken along line 10-10.

FIG. 11 is an enlarged partial cross-sectional view of the syringe assembly of FIG. 1 taken along line 3-3 showing the syringe assembly before use.

FIG. 11A is an enlarged partial cross-sectional view similar to FIG. 11 but rotated 90°.

FIG. 12 is an enlarged partial cross-sectional view of the syringe assembly of FIG. 1 taken along line 3-3 showing the syringe assembly after a first aspiration stroke.

FIG. 12A is an enlarged partial cross-sectional view similar to FIG. 12 but rotated 90°.

FIG. 13 is an enlarged partial cross-sectional view of the syringe assembly of FIG. 1 taken along line 3-3 showing the syringe assembly during a first dispensing stroke.

FIG. 13A is an enlarged partial cross-sectional view similar to FIG. 13 but rotated 90°.

FIG. 14 is an enlarged partial cross-sectional view of the syringe assembly of FIG. 1 taken along line 3-3 showing the syringe assembly at the start of a second aspiration stroke.

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FIG. 14A is an enlarged partial cross-sectional view similar to FIG. 14 but rotated 90°.

FIG. 15 is an enlarged partial cross-sectional view of the syringe assembly of FIG. 1 taken along line 3-3 showing the syringe assembly after a second dispensing stroke.

FIG. 15A is an enlarged partial cross-sectional view similar to FIG. 15 but rotated 90°.

FIG. 16 is an enlarged partial cross-sectional view of the syringe assembly of FIG. 1 taken along line 3-3 showing a position of the internal components in the event of an attempt to withdraw the plunger rod after the second dispensing stroke.

FIG. 16A is an enlarged partial cross-sectional view similar to FIG. 16 but rotated 90°.

FIG. 17 is an enlarged partial cross-sectional view similar to the syringe assembly of FIG. 16 showing the interaction of additional structure to prevent reuse.

FIG. 17A is an enlarged partial cross-sectional view similar to FIG. 17 but rotated 90°.

FIG. 18 is an enlarged partial cross-sectional view similar to the syringe assembly of FIG. 17 showing a discontinuity in the syringe barrel for engaging the locking element.

DETAILED DESCRIPTION OF THE INVENTION

The present invention is directed to a syringe assembly having a passive disabling mechanism. The disabling mechanism enables variable dosages by the syringe assembly and enables a selected number of cycles or strokes by the plunger rod before being automatically disabled. In one preferred embodiment, the disabling mechanism provides two aspirating and two dispensing cycles before being automatically disabled. The assembly enables the aspiration and dispensing of a selected volume of a diluent into a vial to reconstitute a drug, pharmaceutical agent, or other substance and then aspirating the reconstituted substance back into the syringe. A selected volume of the reconstituted substance can be injected or delivered to a patient where the volume of the substance that is delivered can be the same or different than the volume of the substance aspirated into the syringe barrel. The syringe is automatically disabled after the injection or delivery stroke by retracting the plunger rod, which activates the disabling mechanism.

The disabling mechanism is actuated by the axial movement of the plunger rod with respect to the syringe barrel and to the stopper, by moving the plunger rod in the aspirating direction. The stopper is coupled to the plunger rod to allow limited axial movement of the stopper with respect to the plunger rod. The disabling mechanism moves through a series of stages by reversing the direction of the axial movement of the plunger rod with respect to the stopper to move the mechanism in a step-wise manner to the disabling position. The disabling position of the mechanism is attained by the relative movement between the plunger rod and the stopper and is not dependent on the position of the stopper within the syringe barrel or the length of the stroke by the stopper. In this manner, the syringe assembly is able to dispense a desired volume of the drug or other substance, and the disabling mechanism can be actuated after the final dispensing or injection stroke regardless of the position of the stopper in the syringe barrel. By actuating the disabling mechanism, the stopper cannot be retracted to aspirate a substance into the syringe barrel but allows any substance remaining in the syringe barrel to be dispensed.

Referring to the drawings, a syringe assembly 100 having a disabling mechanism includes a syringe barrel 102 and a

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plunger assembly 104. Barrel 102 includes a cylindrical sidewall 106 having an inside surface 107 defining a chamber 109 for retaining fluid, an open proximal end 113 and a distal end 115 including a distal wall 117 having a passageway 119 therethrough in fluid communication with the chamber. In this embodiment, the distal wall of the barrel includes an elongate tip extending distally therefrom and having a passageway in fluid communication with the passageway in the distal wall. In this embodiment barrel 102 also includes a needle cannula 170 having a proximal end 171, a distal end 172 and a lumen 173 therethrough. The proximal end of the needle cannula is attached to elongate tip 103 so that the lumen of the needle cannula is in fluid communication with passageway 119 in the barrel.

Plunger assembly 104 includes an elongate hollow plunger rod 108, a stopper 128 and a locking element 130. Plunger rod 108 includes a proximal end 111, an open distal end 110 and an interior surface 116 and at least one aperture or recess 114 in the interior surface at the distal end of the plunger rod. The recess includes a distal face 121. In this embodiment, there are two recesses 114 having distal faces 121. The interior surface at the distal end of the plunger rod includes at least one detent. In this embodiment the at least one detent on the interior surface of the distal end of the plunger rod includes four axially spaced detents 118 with two detents on each side of the plunger rod. Each pair of detents is shaped to form axially spaced steps 120 with each step having a blunt surface 122 at its distal end extending inwardly from the interior surface of the plunger rod.

Stopper 128 includes a circularly-shaped sealing element 144 having a peripheral surface 145 forming a seal with the inside surface of the barrel. A boss member 134 extends proximally from the sealing element and includes at least one boss detent and in this embodiment, contains two boss detents 136. At least one cantilevered arm extends proximally from the sealing element and in this embodiment there are two cantilever arms 140 extending proximally from the sealing element. Each of the cantilevered arms includes an outwardly extending rib 142. The rib is sized to fit within recess 114 in the plunger rod. The axially spaced boss detents 136 each include an incline surface 137 extending proximally inwardly and a blunt surface 138 at the distal end of each of the inclined surfaces. The stopper is preferably integrally formed of thermoplastic material such as polyethylene. The circularly-shaped sealing element and/or the peripheral sealing surface thereon may be made of elastomeric materials such as thermoplastic elastomers, natural rubber, synthetic rubber and combinations thereof.

Locking element 130 includes a central body portion 148 having an aperture 152 therethrough and at least one cantilevered leg 150 extending distally outwardly from the body portion and at least one finger element 154 extending inwardly from the aperture. In this embodiment, at least two cantilevered legs with each of the cantilevered legs having a sharp free end 155 directed outwardly for engaging the inside surface of the barrel. The configuration of sharp free end 155 can be any configuration capable of engaging the inside surface of the barrel, such as a sharp edge or one or more pointed teeth and the like. The locking element may be made of a variety of materials, or combinations of materials, however, it is preferred to have the sharp free ends made of metal and it is also preferred that the entire locking element be made of integrally formed from sheet metal such as stainless steel.

In this preferred embodiment plunger assembly 104 is assembled by inserting locking element 130 into the distal end of plunger rod 108. Boss 134 of stopper 128 is then

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inserted into the distal end of the plunger rod through aperture 152 of locking element 130 so that cantilevered legs 150 extend toward circularly-shaped sealing element 144 of the stopper as illustrated in FIGS. 9 and 10.

As will be explained in more detail hereinafter, the plunger assembly is then inserted into barrel 102 through open proximal end 113 to the initial position illustrated in FIGS. 11 and 11A. In the initial position of the syringe element, locking element 130 is positioned with its sharp free ends 155 contacting the interior surface of the plunger rod proximally of axially spaced steps 120. Boss member 134 is positioned in aperture 152 of locking element 130 so that finger elements 154 contact boss member proximally of boss detents 136. Outwardly extending ribs 142 of cantilever arms 140 are positioned in recesses 114 in the plunger rod. Ribs 142 are configured to complement the recesses 114 for allowing limited axial movement of the stopper with respect to the plunger rod. The stopper 128 further includes stabilizing member 146 positioned proximally with respect to sealing element 144 and has an outer dimension complementing the other dimension of the sealing element as shown in FIG. 11A, stabilizing member 146 has an outer dimension to contact the inner surface of the syringe barrel and is spaced from sealing element 144 to assist in stabilizing stopper 128 to maintain the stopper and boss member 134 in an orientation substantially parallel to the axis of the syringe barrel. In the position illustrated in FIGS. 11 and 11A, syringe assembly 100 is ready to use for drawing liquid into the chamber of the barrel.

As will now be shown, the operation of the plunger assembly of this embodiment includes a first aspiration stroke followed by a first dispensing stroke, a second aspiration stroke and a final dispensing stroke after which the syringe is disabled. The disabling elements prevent or inhibit movement of stopper 128 in a proximal aspirating direction thereby limiting the function of the syringe assembly to a single use. The maximum number of strokes being limited by a number of axially positioned detents in the plunger rod and the number of axially positioned boss detents on the stopper. However, the actual number of strokes the syringe may make will be determined by the position of the locking element with respect to the detents in the plunger rod and the detents on the stopper at the time of first use. For example, a syringe with two plunger detents and two stopper detents can be supplied to the end user as a syringe capable of two strokes or four strokes. This is an important feature of the present invention since a single syringe assembly can be provided with different stroke limitations before disabling.

The syringe assembly may now be used to draw liquid, such as a sterile water diluent into the chamber of the barrel by applying a proximally directed force to a thumb press 123 on the proximal end of the plunger rod while holding the syringe barrel. As illustrated in FIGS. 12 and 12A, this causes the plunger rod to move proximally with respect to the stopper until the free end of cantilevered legs 150 moves distally along inner surface 116 of the plunger rod and snaps past blunt surface 122 of the proximal most axially spaced steps 120, as best illustrated in FIG. 12. Also, during this first aspiration stroke outwardly extending ribs 142 engage distal surface 121 of the recesses 114 in the plunger rod as best illustrated in FIG. 12A. When ribs 142 engage distal surface 121 the stopper is drawn proximally with respect to the barrel as the plunger rod moves. The stopper is now moved proximally, through action of the plunger rod, until the desired volume of liquid is in the chamber as determined by the user.

The liquid diluent in the chamber may now be discharged into a vial of dry medication such as lyophilized medication, for reconstitution. This first dispensing stroke is accomplished by moving the plunger rod in a distal direction while holding the barrel. A barrel flange **124** is provided on the proximal end of the barrel to help control motion of the barrel during use of the syringe assembly. As best illustrated in FIGS. **13** and **13A**, as the plunger rod moves distally, locking element **130** moves with the plunger rod dragging the locking element with it so that finger elements **154** on the locking element slide from the proximal most to the distal most boss detent by riding up inclined surface **137** and falling into the second detent. When the plunger rod contacts the stopper, the stopper will begin moving in a distal direction along with the plunger rod to discharge liquid diluent from the chamber into, for example, a vial of lyophilized medication.

When the diluent and the lyophilized medication are mixed the syringe assembly of the present invention may now be used to withdraw the reconstituted, ready-to-inject medication into the chamber of the syringe barrel, as best illustrated in FIGS. **14** and **14A**, by applying a proximally directed force to the plunger rod while holding the syringe barrel. Proximally directed force will cause the plunger rod to move in a proximal direction while locking element **130** will remain relatively stationary due to its connection to the boss detent on the stopper. Proximal motion of the plunger causes the locking element to move distally along the inside surface of the plunger rod so that the sharp free end **155** of the cantilever legs moves from the proximal-most axially-spaced steps **120** in the plunger rod to the second more distal axially-spaced steps **120**. Proximal motion of the plunger rod also causes outwardly extending ribs **142** of cantilever arms **140** to engage distal surfaces **121** of recesses **114** in the plunger rod so that the stopper now moves proximally with the plunger rod drawing the reconstituted medication into the chamber of the syringe barrel to an amount determined by the user. An advantage of the present invention is that the amount of medication drawn into the chamber, and therefore the maximum amount of medication that can be delivered, is determined by the user at the time of use and not by the placement of the components at the time of manufacture.

The syringe assembly of the present invention is now ready for a second and final dispensing stroke which is best illustrated in FIGS. **15-15A**. The medication is delivered to the patient by applying a distally directed force to the plunger rod causing the plunger rod to move in a distal direction with respect to the barrel. As the plunger rod advances in a distal direction the engagement of sharp free ends **155** of the locking element in with the distal-most blunt surfaces **122** of axially-spaced steps **120** moves the locking element distally so that finger elements **154** of the locking element ride over the distal-most inclined surface **137** of the boss detents distally past the most distal boss detent **136**. When the distally moving plunger rod contacts the stopper, both the stopper and the plunger rod move toward the distal end of the barrel to discharge the contents of the chamber through the passageway.

The syringe assembly has now been used and is ready to be discarded. Any attempt to move the plunger rod in a proximal direction to refill the syringe assembly for further use will cause the locking element to disable the syringe. Specifically, as best illustrated in FIGS. **16** and **16A**, moving the plunger rod in a proximal direction will allow the plunger rod to move a short distance until the sharp free ends **155** of the locking clip snap past the end of the plunger rod and engage the inside surface **107** of the barrel side wall.

Further proximal motion of the plunger will be resisted by the locking element's engagement to the inside surface of the barrel sidewall. In addition, as illustrated in FIG. **17**, cam surface **125** on the stopper is positioned to force sharp free ends **155** further into the syringe barrel wall as more proximally directed force is used in an attempt to improperly reuse the syringe. Accordingly, increased force to pull the plunger rod out of the syringe barrel results in increased force of engagement of the sharp free ends of the locking element into the barrel. It is desirable to provide a cut-out area **126** in the distal end of the plunger rod along the path of the sharp free ends of the locking element for supporting the locking element and allowing it to engage the inside surface of the barrel. Further, the area at the end of the plunger rod on the area around the cutout can be configured to support the locking element so that if the user accidentally withdraws the plunger rod a second time before delivering the final dose of medication the medication may still be delivered even though the locking element sharp free ends are touching the barrel so long as they are moved in a distal direction and urged not to engage the inside surface of the barrel by the cut-out area and the plunger rod. In this case the area around the cutout supports and limits the motion and helps prevent deformation of the sharp free end of the locking element.

It is also within the purview of the present invention to provide a discontinuity such as a recess or projection on the interior surface of the barrel, as illustrated in FIG. **18**, to further improve the engagement of the sharp free end of the locking element with the interior surface of the barrel. In FIG. **18** syringe barrel **102** includes a discontinuity in the form of an inwardly directed projection **127** on inside surface **107** of the barrel. In this embodiment, projection **127** is an annular ring projecting into the barrel and extending 360° around the inside surface. The discontinuity may be in the form of an annular projection, an annular recess or one or more projections or recesses shaped to engage locking element, all positioned within the barrel to engage sharp free end **155** of locking element **130** to further increase the grip of the locking element on the barrel and inside surface.

The present syringe assembly provides an improvement over prior art devices by allowing a variable dose of diluent, chosen by the user at the time of use, to be drawn into the syringe, dispensing the diluent into a vial containing a substance to be reconstituted, drawing a selected amount of the reconstituted substance back into the syringe and then delivering the contents of the syringe. The selected amount of the reconstituted substance may be equal or less than the full volume reconstituted at the discretion of the user. The syringe assembly is automatically disabled after the final injection stroke by reversing the direction of the movement of the plunger rod from the dispensing direction to the aspirating direction. After the injection stroke of the syringe plunger the plunger rod is retracted to activate the disabling mechanism to prevent axial movement of the stopper toward the proximal end of the syringe barrel thereby preventing the stopper from being removed and preventing reuse of the syringe.

When the present syringe assembly has two or more detents on the stopper and in the plunger rod, the maximum number of strokes the syringe assembly will allow can be varied by the initial position of the locking element with respect to the stopper detents and the plunger rod detents.

While various embodiments have been chosen to illustrate the invention, it will be appreciated that changes and modifications can be made without departing from the scope of the invention.

What is claimed is:

1. An operable syringe assembly having passive disabling structure comprising:

a barrel including a cylindrical side wall having an inside surface defining a chamber for retaining fluid, an open proximal end and a distal end including a distal wall having a passageway therethrough in fluid communication with said chamber;

an elongate hollow plunger rod having a proximal end, an open distal end and an interior surface, at least one recess in said interior surface at said distal end of said plunger rod, said recess having a distal face, at least one detent on said interior surface at said distal end of said plunger rod;

a stopper including a circular shaped sealing element having a peripheral surface forming a seal with said inside surface of said barrel, a boss member projecting proximally from said sealing element, at least one boss detent on said boss, at least one cantilevered arm extending proximally from said sealing element, said at least one cantilevered arm having an outwardly extending rib near its free end, said rib being sized to fit within said recess in said plunger rod;

a locking element including a central body portion having an aperture therethrough, at least one cantilevered leg extending distally outwardly from said body portion, and at least one finger element extending proximally inwardly from said aperture, said at least one leg having a sharp free end directed outwardly for engaging said inside surface of said barrel;

said locking element being positioned with said sharp free end contacting said interior surface of said plunger rod proximally of said at least one detent, said boss member being positioned in said aperture of said locking element wherein said at least one finger element is contacting said boss proximally of said at least one boss detent, and said outwardly extending rib being positioned in said recess of said plunger rod, so that applying a proximally directed force to said plunger rod while holding said barrel causes said plunger rod to move proximally with respect to said stopper until said free end of said cantilevered leg moves distally along said inner surface of said plunger rod to said at least one detent and said outwardly extending rib on said cantilevered arm engages said distal face of said recess to move said stopper in a proximal direction for a selected distance, and subsequently applying a distally directed force to said plunger rod to discharge fluid from said chamber through said passageway causes said plunger rod to move in a distal direction along with said locking element due to its engagement with said at least one detent until said finger element of said locking element rides over said at least one boss detent and into contact with said at least one boss detent and said plunger rod contacts said stopper to move said stopper distally to discharge fluid from said chamber, after which applying a proximally directed force to said plunger rod will cause said plunger rod to move proximally until said free end of said cantilevered leg moves distally along said inner surface past said distal end of said plunger rod so that said sharp end of said at least one cantilevered leg engage said inside of said barrel to help prevent proximal movement of said stopper to render said syringe assembly unusable.

2. The syringe assembly of claim 1 wherein said at least one detent in said plunger rod includes two axially spaced detents and said at least one detent on said boss includes two

axially spaced boss detents so that said plunger rod can be moved distally two times before proximal motion of said plunger rod causes said locking element to engage said inside surface of said barrel.

3. The syringe assembly of claim 2 wherein said two axially spaced detents in said plunger rod include two axially spaced steps each having a blunt surface at its distal end extending inwardly from said interior surface.

4. The syringe assembly of claim 2 wherein said two axially spaced boss detents each include an inclined surface extending proximally inwardly and a blunt surface at a distal end of each of said inclined surfaces extending radially inwardly.

5. The syringe assembly of claim 2 said at least one cantilevered leg of said locking element includes two cantilevered legs positioned on opposite sides of said central body portion and two additional axially spaced detents are provided in said plunger rod opposite said two axially spaced detents.

6. The syringe assembly of claim 5 further including two radial projections on said stopper positioned to engage and force said two cantilevered legs outwardly when excessive proximally directed force is applied to said plunger rod in an attempt to overcome said locking element's engagement of said inside surface of said barrel.

7. The syringe assembly of claim 1 wherein said at least one cantilevered arm of said stopper includes two cantilevered arms positioned on opposite sides of said boss and said at least one recess in said inner surface of said plunger rod includes two recesses positioned on opposite sides of said plunger rod and positioned to receive said outward extending ribs of said cantilevered arms.

8. The syringe assembly of claim 1 having a discontinuity on said inside surface of said barrel sidewall positioned to engage said sharp free end of said locking element when said sharp free end is contacting said inside surface of said barrel.

9. The syringe assembly of claim 1 wherein said distal wall of said barrel further includes an elongate tip extending distally therefrom having a passageway in fluid communication with said passageway in said distal wall.

10. The syringe assembly of claim 1 further including a needle cannula having a distal end, a proximal end and a lumen therethrough, said proximal end of said needle cannula being connected to said distal end of said barrel so that said lumen is in fluid communication with said passageway.

11. The syringe assembly of claim 1 wherein said locking element is made of sheet metal.

12. The syringe assembly of claim 1 wherein said stopper is integrally formed of thermoplastic material.

13. An operable syringe assembly having passive disabling structure comprising:

a barrel including a cylindrical side wall having an inside surface defining a chamber for retaining fluid, an open proximal end and a distal end including a distal wall having an elongate tip extending distally therefrom and a passageway therethrough in fluid communication with said chamber;

an elongate hollow plunger rod having a proximal end, an open distal end and an interior surface, said interior surface at said distal end of said plunger rod including two opposed recesses and two pairs of axially spaced opposed steps, each step extending inwardly from said sidewall and having a blunt surface at its distal end and each recess having a distal face;

a stopper including a circularly shaped sealing element having a peripheral surface forming a seal within said inside surface of said barrel, a boss member projecting

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proximally from said sealing element, two axially spaced detents on said boss each having an inclined surface extending proximally inwardly and a blunt surface extending radially away from said boss, two cantilevered arms extending proximally from said seal-

ing element, each cantilevered arm having an outwardly extending rib near its free end, said rib being sized to fit within said recesses in said plunger rod;

a locking element including a central body portion having an aperture therethrough and two opposed cantilevered legs extending distally outwardly from said body portion, two finger elements extending proximally inwardly from said aperture, said legs having free ends including a hook-shaped portion, said hook-shaped portion having a sharp end directed proximally outwardly from engaging said inside surface of said barrel and said blunt surface in said plunger rod;

said locking element being positioned with said sharp ends of said hook-shaped portions contacting said interior surface of said plunger rod proximally of said blunt surface of said steps, said boss member being positioned in said aperture of said locking element wherein said finger elements contact said boss proximally of said blunt surface of said detents, and said outwardly extending ribs being positioned in said recess of said plunger rod, so that applying a proximally directed force to said plunger rod while holding said barrel causes said plunger rod to move proximally with respect to said stopper until said hook-shaped portions of said cantilevered legs move distally along said inner surface of said plunger rod and moves outwardly at said blunt surface of proximal-most of said two steps and said outwardly extending ribs on said cantilevered arms engage said distal face of said recesses to move said stopper in a proximal direction for a selected distance, and subsequently applying a distally directed force to discharge fluid from said chamber through said passageway causes said plunger to move in a distal direction with said locking element due to said locking element's engagement with the proximal-most of said blunt surfaces until said finger elements of said locking element ride over said inclined surface of the proximal-most said two detents and into contact with said blunt surface and said plunger rod contacts said stopper to move said stopper distally to discharge fluid from said chamber, an additional proximal and distal movement of said plunger rod to draw fluid into said chamber and to deliver fluid from said chamber will cause said locking element to be repositioned so that said hook-shaped portions of said legs engage the distal-most of said two steps and said finger elements engage the distal-most of said blunt surfaces, after which applying a proximally directed force to said plunger will cause said plunger rod to move proximally until said hook-shaped portions of said cantilevered legs move distally along said inner surface of said plunger rod past said distal end of said plunger rod so that said sharp ends of said hook-shaped projections engage said inside surface of said barrel to help prevent proximal motion of said stopper to render said syringe assembly unusable.

14. The syringe assembly of claim **13** wherein said two axially spaced boss detents each include an inclined surface extending proximally inwardly and a blunt surface at a distal end of each of said inclined surfaces extending radially inwardly.

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15. An operable syringe assembly having passive disabling structure comprising:

a barrel including a cylindrical sidewall having an inside surface defining a chamber for retaining fluid, an open proximal end and a distal end including a distal wall having a passageway therethrough in fluid communication with said chamber;

an elongate hollow plunger rod having a proximal end, an open distal end and an interior surface;

a stopper including a circular-shaped sealing element having a peripheral surface forming a seal with said inside surface of said barrel and a boss member projecting proximally from said sealing element;

a locking element including a central body portion having at least one cantilevered leg extending distally outwardly from said body portion, said at least one leg having a sharp free end directed outwardly for engaging said inside surface of said barrel, said locking element movably connected to said boss and said locking element movably connected to said plunger rod interior surface; and

means for indexing said locking element distally in said plunger rod during proximal motion of said plunger rod to draw fluid into said chamber and for indexing said locking element distally on said boss of said stopper during distally directed motion of said plunger rod for delivering fluid from said chamber through said passageway and means for engaging said locking element with said inside surface of said barrel sidewall for preventing reuse of said syringe assembly.

16. The syringe assembly of claim **15** wherein said at least one detent in said plunger rod includes two axially spaced detents and said at least one detent on said boss includes two axially spaced boss detents so that said plunger rod can be moved distally two times before proximal motion of said plunger rod causes said locking element to engage said inside surface of said barrel.

17. The syringe assembly of claim **16** wherein said two axially spaced detents in said plunger rod include two axially spaced steps each having a blunt surface at its distal end extending inwardly from said interior surface.

18. The syringe assembly of claim **15** said at least one cantilevered leg of said locking element includes two cantilevered legs positioned on opposite sides of said central body portion and said at least one detent in said interior surface of said distal end of said plunger rod includes two detents positioned on opposite side of said plunger rod.

19. The syringe assembly of claim **18** further including two radial cam projections on said stopper positioned to contact and force said two cantilevered legs outwardly when excessive proximally directed force is applied to said plunger rod in an attempt to overcome said locking element's engagement to said inside surface of said barrel.

20. The syringe assembly of claim **15** wherein said means for engaging said locking element with said inside surface of said barrel side wall includes an opening in said distal end of said plunger rod to shorten the axially length of the interior surface in the area of the opening and to allow said sharp free end to project outwardly past the plunger rod and into the inside surface of the barrel.

21. The syringe assembly of claim **15** wherein means for indexing includes at least one detent on said interior surface of said distal end of said plunger rod, at least one boss detent on said boss, at least one cantilevered arm on said stopper having an outwardly extending rib near its free end sized to engage a recess in inside surface of said plunger rod and said locking element including at least one finger element

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extending inwardly from an aperture in said central body portion, said locking element being positioned with said sharp free end contacting said interior surface of said plunger rod proximally of said at least one detent, said boss member being positioned in said aperture of said locking element wherein said at least one finger element is contacting said boss proximally of said at least one boss detent and said outwardly extending rib being positioned in said recess of said plunger rod.

22. The syringe assembly of claim 15 wherein said at least one cantilevered arm of said stopper includes two cantilevered arms positioned on opposite sides of said boss and said at least one recess in said inner surface of said plunger rod includes two recesses positioned on opposite sides of said plunger rod and positioned to receive said outward extending ribs of said cantilevered arms.

23. The syringe assembly of claim 15 having a discontinuity on said inside surface of said barrel sidewall posi-

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tioned to engage said sharp free end of said locking element when said sharp free end is contacting said inside surface of said barrel.

24. The syringe assembly of claim 15 wherein said distal wall of said barrel further includes an elongate tip extending distally therefrom having a passageway in fluid communication with said passageway in said distal wall.

25. The syringe assembly of claim 15 further including a needle cannula having a distal end, a proximal end and a lumen therethrough, said proximal end of said needle cannula being connected to said distal end of said barrel so that said lumen is in fluid communication with said passageway.

26. The syringe assembly of claim 15 wherein said locking element is made of sheet metal.

27. The syringe assembly of claim 15 wherein said stopper is integrally formed of thermoplastic material.

* * * * *

UNITED STATES PATENT AND TRADEMARK OFFICE
CERTIFICATE OF CORRECTION

PATENT NO. : 7,331,934 B2
APPLICATION NO. : 10/838687
DATED : February 19, 2008
INVENTOR(S) : Suresh et al.

Page 1 of 1

It is certified that error appears in the above-identified patent and that said Letters Patent is hereby corrected as shown below:

Title Page:

Insert Item -- (73 Assignee: Becton, Dickinson and Company, Franklin Lakes, NJ --.

Signed and Sealed this

Twenty-eighth Day of October, 2008

A handwritten signature in black ink that reads "Jon W. Dudas". The signature is written in a cursive style with a large, looped initial "J".

JON W. DUDAS

Director of the United States Patent and Trademark Office